

# Artificial Intelligence for Osteoporosis Diagnosis, Risk Prediction and Therapy: Current Advances, Clinical Challenges, and Future Perspectives

Zhaochen Zhang<sup>1,\*</sup>, Yuxi He<sup>2,\*</sup>, Zhanhao Mo<sup>3,\*</sup>, Peng Zhang<sup>4</sup>, Zhenya Tian<sup>5</sup>, Lanfeng Huang<sup>1</sup>

<sup>1</sup>Department of Orthopedics, The Second Hospital of Jilin University, Changchun, Jilin, People's Republic of China; <sup>2</sup>Department of Ophthalmology, The Second Hospital of Jilin University, Changchun, Jilin, People's Republic of China; <sup>3</sup>Department of Radiology, China-Japan Union Hospital of Jilin University, Changchun, Jilin, People's Republic of China; <sup>4</sup>Department of Radiology, The Second Hospital of Jilin University, Changchun, Jilin, People's Republic of China; <sup>5</sup>The First Norman Bethune Clinical Medical College, Jilin University, Changchun, Jilin, People's Republic of China

\*These authors contributed equally to this work

Correspondence: Lanfeng Huang, Department of Orthopedics, The Second Hospital of Jilin University, Changchun, Jilin, People's Republic of China, Email [hlf@jlu.edu.cn](mailto:hlf@jlu.edu.cn)

**Abstract:** Osteoporosis (OP) is a chronic systemic skeletal disorder that predominantly affects the elderly. It is characterized by an imbalance in bone homeostasis, reduced bone mass, microarchitectural deterioration of bone tissue, and increased bone fragility, ultimately leading to a higher risk of fractures and related complications. With the progression of global population aging, the prevalence of OP continues to rise, underscoring the importance of early diagnosis and timely intervention. However, the diagnosis and management of OP—particularly its early detection—remain limited by material constraints such as diagnostic equipment and by subjective factors including clinician experience, which hinder widespread screening. In recent years, artificial intelligence (AI) has emerged as a transformative technology with advantages of efficiency, objectivity, and scalability, and has been increasingly integrated into various medical domains. For example, AI-assisted musculoskeletal measurements on leg and foot radiographs can reduce the measurement time from 166 seconds to 40 seconds, resulting in an overall efficiency improvement of approximately 70%. Applying AI to the diagnosis and treatment of OP can reduce human error, save labor costs, and improve diagnostic accuracy and clinical efficiency. Numerous studies have investigated AI-based approaches in OP-related research and clinical practice. Despite these promising developments, several important limitations should be acknowledged. Considerable heterogeneity exists among published studies regarding patient populations, AI algorithms, and evaluation metrics. Besides, consistent external validation remains insufficient in many studies. Challenges related to data imbalance and potential selection bias further highlight the need for standardized reporting frameworks and multicenter collaborative research to promote safe clinical adoption of AI technologies in osteoporosis management. This review summarizes current AI applications in OP diagnosis, risk prediction and therapy. We highlight key methodological limitations and emerging trends, aiming to guide future research and facilitate safe clinical implementation of AI in OP management.

**Keywords:** artificial intelligence, osteoporosis, image recognition, big data, bioinformatics

## Introduction

OP is a chronic systemic skeletal disorder characterized by an imbalance in bone homeostasis, decreased bone mass, alterations in trabecular microarchitecture, and increased bone fragility.<sup>1</sup> Epidemiological data indicate that in China alone, approximately 49 million women and 22.8 million men aged over 50 years are affected by OP, highlighting a substantial disease burden in the world's largest aging population.<sup>2</sup> Similarly, in the United States, musculoskeletal disorders affect more than 121 million individuals and remain one of the leading causes of disability across all disease categories.<sup>3</sup> It primarily affects elderly individuals, particularly postmenopausal women. According to data from the International Osteoporosis Foundation (IOF), one in three women over the age of 50 worldwide are affected by OP, and

many eventually develop serious complications, the most prominent of which is osteoporotic fracture (OF).<sup>4,5</sup> Globally, OP is estimated to cause 8.9 million fractures annually.<sup>6</sup> With the acceleration of population aging, improving early identification and management of OP has become an urgent public health priority.

Currently, the diagnosis of OP mainly relies on dual-energy X-ray absorptiometry (DXA). When a patient's bone mineral density (BMD) is equal to or less than 2.5 standard deviations below the mean value for healthy young adults, OP can be diagnosed.<sup>7,8</sup> However, both the availability and utilization rate of DXA remain relatively low, and patient adherence to DXA screening is often poor,<sup>9</sup> creating a significant barrier to early diagnosis. On the treatment side, OP management still depends heavily on traditional drug development, which is time-consuming, costly, and has a high failure rate, thereby increasing the difficulty of effective intervention.<sup>10</sup>

AI—which is rapidly advanced in the past three decades<sup>11</sup>—has been increasingly integrated into diverse areas of medicine,<sup>12</sup> including automated medical image recognition,<sup>13</sup> early disease diagnosis,<sup>14</sup> and drug discovery.<sup>10</sup> Key enabling technologies include machine learning (ML), deep learning (DL),<sup>15</sup> large-scale foundation models,<sup>10</sup> and Generative AI.<sup>16</sup> In recent years, with the development of big data analytics, the convergence of AI and large-scale biomedical data has opened new directions for medical research and clinical applications.<sup>17</sup>

The unique capabilities of AI provide promising solutions to the challenges faced in OP diagnosis and treatment. In diagnostics, AI enables identification of osteoporotic features from common imaging modalities such as X-ray,<sup>18</sup> computed tomography (CT),<sup>19</sup> magnetic resonance imaging (MRI),<sup>20</sup> and ultrasound,<sup>21</sup> offering potential for opportunistic screening. Moreover, AI can identify risk factors and predict the likelihood of OP and OF occurrence.<sup>22</sup> In therapeutics, AI contributes to the discovery of key genetic targets for OP,<sup>23</sup> supports drug development,<sup>10</sup> and predicts clinical outcomes.<sup>24</sup> Despite these promising developments, growing evidence indicates substantial variability in patient populations, algorithm selection, and evaluation strategies, which may limit the comparability, reproducibility, and clinical translation of current AI models.<sup>10,25–27</sup>

Therefore, a comprehensive review that not only summarizes existing applications but also critically examines methodological limitations, validation challenges, and barriers to real-world implementation is needed. This review aims to summarize current applications of AI in OP screening, diagnosis, risk prediction, and treatment research, while critically discussing methodological challenges and future directions to support safe and effective clinical adoption.

## Methods

This narrative review aimed to summarize current applications of AI in osteoporosis OP research and clinical practice. Relevant literature published in English was retrieved from PubMed and IEEE Xplore databases up to May 2026.

In PubMed, both Medical Subject Headings (MeSH) terms and free-text keywords were used. The primary MeSH terms included “Artificial Intelligence” and “Osteoporosis”. Additional keyword searches were also performed using terms such as “machine learning”, “deep learning”, “radiomics” and “osteoporotic fracture” to identify studies related to AI-assisted diagnosis, prediction, and treatment of OP. Furthermore, broader searches using the keyword “artificial intelligence” were conducted to support the introductory overview of AI technologies and concepts.

In IEEE Xplore, literature searches were performed using combinations of keywords including “artificial intelligence and medicine”, “artificial intelligence and orthopedics”, and “artificial intelligence and osteoporosis” to identify engineering- and technology-oriented studies relevant to musculoskeletal medicine and OP research.

Original research articles, reviews, and clinically relevant studies focusing on AI applications in OP diagnosis, prediction, molecular analysis, imaging, or treatment were included. References from relevant articles were also manually screened to identify additional studies of relevance.

## Overview of AI Technologies in Osteoporosis

With the rapid advancement of modern technology, AI has evolved into several major subfields, including ML, DL, natural language processing (NLP), and robotics and automation.<sup>28</sup> Each of these branches has found distinct yet complementary applications in medicine. For instance, ML is widely used for feature extraction and patient data analysis;<sup>29</sup> DL has become fundamental in medical image recognition and classification;<sup>30</sup> NLP assists clinical decision-making by supporting treatment planning, providing alerts, and monitoring adverse events,<sup>31</sup> while robotics and

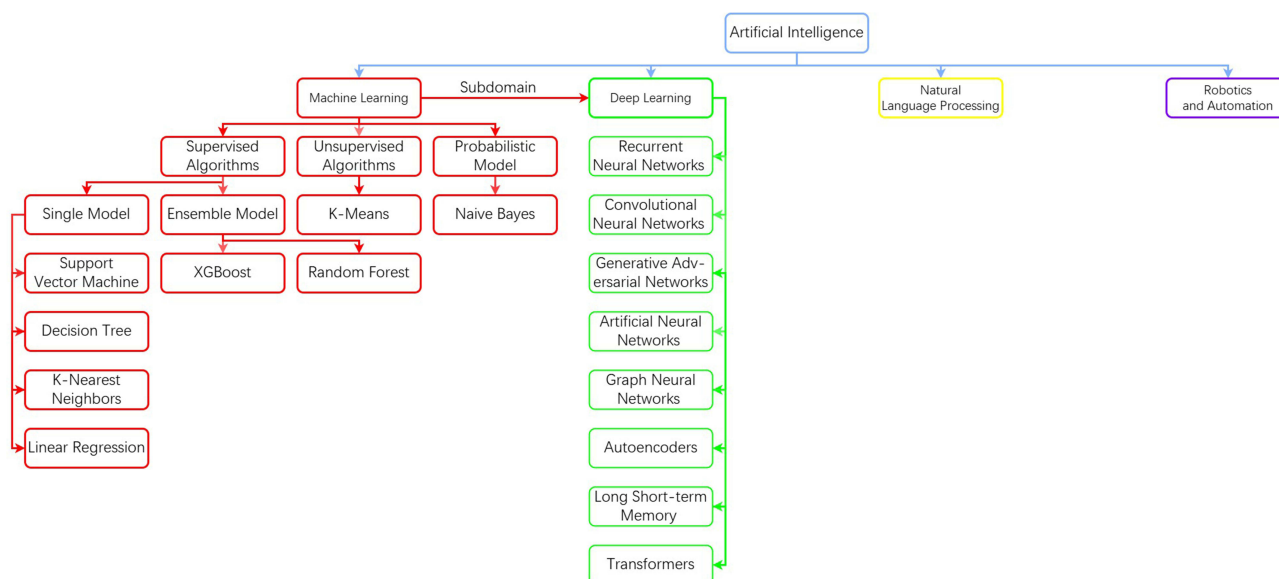
automation are primarily applied in surgical assistance<sup>32</sup> and patient rehabilitation.<sup>33</sup> Although these AI subfields differ in their core methodologies and application domains, they are often interrelated and integrated in clinical research and practice. The interconnections among these branches are illustrated in Figure 1.

## Machine Learning for Osteoporosis Data Analysis

ML is a fundamental branch of AI that has been widely applied in healthcare.<sup>28</sup> It is good at the extraction and interpretation of complex data patterns, particularly for image feature extraction,<sup>40</sup> risk factor assessment,<sup>22</sup> and bioinformatics-based prediction.<sup>23</sup> OP-related data are typically characterized by high dimensionality, heterogeneous data sources (eg., imaging, clinical indicators, and molecular data), and relatively limited sample sizes, which makes ML particularly suitable for OP research. Now ML plays a critical role in disease diagnosis and prognosis, drug discovery and design, gene and molecular mechanism analysis, and risk factor prediction.<sup>41–44</sup>

### ML for Diagnosis and Prognosis of Osteoporosis

Among all the directions, the use of ML for OP diagnosis and prognosis prediction has attracted the most attention. Because in X-ray imaging, manual measurement of OP-related indicators requires substantial manpower and time, which results in low reproducibility of manual classification methods. ML can extract quantitative features such as size, shape, and texture heterogeneity from segmented images, thereby overcoming the above limitations.<sup>45,46</sup> This line of research mainly follows three approaches: image recognition,<sup>47</sup> radiomics,<sup>48</sup> and bioinformatics.<sup>43</sup> Image recognition-based diagnosis and prognosis prediction of OP currently represents one of the most extensively studied directions in ML research. For example, Galbusera F et al applied ML to MRI and X-ray images of the lumbar spine for OP detection. Their study found that two models—CatBoost and Gradient Boosting Classifier (GBC)—achieved the best performance, with CatBoost showing optimal accuracy for MRI (0.90) and GBC for X-rays (0.88).<sup>49</sup> In radiomics-based studies, Zhang et al integrated clinical variables such as CT values and cross-sectional areas (CSA) of the L1 vertebral body, psoas major (PM), and paraspinal muscles (PVM), along with patient demographics, DXA T-scores, and follow-up data. Using a Support Vector Machine (SVM) algorithm, they constructed a multimodal model that combined BMD, CT features, and radiomic-DL signatures to predict OF risk. Furthermore,<sup>40</sup> in bioinformatics-driven analyses, Hu et al identified



**Figure 1** Relationships among commonly used AI technologies.<sup>1,28,34–39</sup> Blue boxes and arrows AI and its relationships with related domains. Red boxes and arrows indicate ML and its subfields, while green boxes and arrows denote DL and its associated areas. Yellow boxes represent NLP, and purple boxes represent robotics and automation. AI mainly consists of four subfields: ML, DL, NLP, and robotics and automation. ML primarily includes three types of algorithms: supervised algorithms, unsupervised algorithms (eg., K-means), and probabilistic models (eg., Naive Bayes). Supervised algorithms can be further divided into single models (eg., LR, SVM, k-NN, and decision trees) and ensemble models (eg., XGBoost and RF). DL is a subfield of ML; however, it has increasingly evolved into a relatively independent research area parallel to traditional machine learning. Its main categories include CNN, RNNs, generative adversarial networks (GANs), ANNs, graph neural networks (GNNs), autoencoders, LSTMs, and transformers.

differentially expressed genes (DEGs) associated with OP through bioinformatic screening and subsequently built an SVM classification model capable of accurately distinguishing osteoporotic samples.<sup>50</sup>

### ML in Molecular Mechanism Research

In addition to imaging analysis, ML has also been increasingly applied to molecular mechanism exploration in OP research. By integrating bioinformatics techniques with ML algorithms, researchers can identify disease-related genes, signaling pathways, and molecular biomarkers associated with OP progression. For instance, Zhang et al investigated the molecular pathways underlying plasticizer-induced OP by employing ten ML algorithms (including Lasso, SVM, and Random Forest [RF]) to build 113 predictive models, ultimately identifying six core genes: CKM, TACR3, SOAT2, ERAP2, SGK1, and MMP12.<sup>51</sup> Similarly, Yang et al examined the genetic correlation between chronic hepatitis B virus (HBV) infection and OP. Using LASSO regression, recursive feature elimination (RFE), and three ML models (RF, SVM, and Gradient Boosting Machine [GBM]), they screened for disease-associated genes and identified 18 key genes, including USP10, ERAL1, and ECM1.<sup>52</sup> These approaches demonstrate the potential of ML to improve the efficiency and precision of molecular-level investigations. Detailed applications of AI in genetic and molecular studies of OP are further discussed in [Artificial Intelligence in Genetic and Molecular Studies of Osteoporosis](#).

### ML for Risk Factor Prediction

Risk factor prediction represents another major application area of ML in OP research. Chang et al compared the predictive accuracy of ML algorithms with multiple linear regression (MLR) for changes in BMD ( $\Delta T$ -score) among 1,698 postmenopausal women. By incorporating biochemical and lifestyle variables, and using RF, XGBoost, Naïve Bayes (NB), and Stochastic Gradient Boosting (SGB) algorithms, they identified key determinants such as educational level that significantly influenced bone density variation.<sup>53</sup> Similarly, Xu et al integrated demographic variables and blood-based biomarkers into ML models, demonstrating that serum biomarkers can effectively distinguish individuals with low BMD, thereby highlighting their potential as noninvasive diagnostic indicators.<sup>54</sup>

### ML for Drug Design in Osteoporosis

Although studies applying ML to drug design for OP remain limited, emerging work has demonstrated promising results. Yang et al combined LASSO, SVM-RFE, and RF algorithms with bioinformatics analyses to identify shared diagnostic genes for comorbid OP and sarcopenia, including CHST3, PGBD5, and SLIT2. These genes were subsequently queried in the Connectivity Map (CMap) database to predict potential therapeutic compounds, such as PU-H71, Scandenin, and BMS-345541.<sup>55</sup>

In summary, ML has shown broad and expanding applications in OP research, encompassing diagnostic imaging, molecular biology, pharmacology, and clinical risk assessment. With ongoing advances in data integration and algorithmic modeling, ML-based approaches are expected to play an increasingly significant role in future OP studies.

## Deep Learning in Osteoporosis Imaging

DL is a subfield of ML that enables computational models composed of multiple processing layers to learn data representations at various levels of abstraction. DL models demonstrating remarkable capability in feature learning.<sup>34</sup> In the medical domain, DL has been extensively applied to the analysis and interpretation of medical imaging data.<sup>30</sup> So DL is particularly well suited for OP -related imaging tasks, as it enables automatic extraction of hierarchical features from raw radiographic and CT images. These models can capture complex trabecular and cortical bone patterns that are difficult to quantify manually, reducing reliance on handcrafted features and improving robustness across different imaging conditions.<sup>46</sup>

In the context of OP, DL has primarily been used for image-based diagnosis and auxiliary detection.<sup>56</sup> Large-scale application of DL models can significantly reduce manual workload, shorten image reading time, and minimize the impact of human subjectivity. For example, Ho et al developed a DL model named DeepDXA-Hand, based on the efficient convolutional neural network (CNN) architecture HarDNet, to detect OP noninvasively using hand X-rays. The model achieved high diagnostic performance, with a sensitivity of 0.73, specificity of 0.83, and accuracy of 0.80.<sup>57</sup> Similarly, Pan et al constructed a DL-based segmentation framework that integrated multiple radiomic features to classify OP from chest CT scans. Their model achieved outstanding area under the curve (AUC) values of 0.992 for normal bone density, 0.973 for osteopenia, and 0.989 for OP, demonstrating exceptional diagnostic capability.<sup>58</sup>

Beyond imaging applications, DL has also been explored in drug discovery and optimization related to OP. According to Xu et al, DL techniques can facilitate drug target identification, lead compound screening and optimization, prediction of physicochemical properties, drug–drug interaction modeling, and synthetic route design. Compared with traditional drug discovery methods—which are often costly, time-consuming, and have low success rates, DL-based approaches offer higher efficiency and scalability. However, specific studies applying DL to OP drug development remain limited, suggesting that this may become a promising direction for future research.<sup>10</sup>

## Potential Roles of Other Models in Osteoporosis Research

In addition to traditional ML and DL approaches, large models have recently emerged as one of the most actively studied branches of AI. Current research on the application of large models in OP primarily focuses on evaluating their potential utility in OP-related drug development.<sup>10</sup> Although no concrete clinical or laboratory studies have yet been conducted, the intrinsic advantages of large models—such as their ability to process high-dimensional data, perform multitask learning<sup>10</sup>—suggest significant potential for future applications in OP research. In OP research and clinical practice, large language models may play a supportive role by processing unstructured clinical narratives, guideline documents, and electronic health records(EHRs).<sup>59</sup>

While AI encompasses a wide range of techniques and methodologies, its most mature and clinically impactful applications in OP to date have been concentrated in disease detection and diagnosis. In particular, advances in ML and DL have enabled the extraction of subtle skeletal features from routine medical images, laying the foundation for AI-assisted imaging-based diagnosis.<sup>60</sup>

## AI in Osteoporosis Diagnosis

With the progression of population aging, the incidence of OP has been increasing steadily each year, making early case identification particularly critical. Although DXA remains the gold standard for diagnosing OP, its limited accessibility and low screening coverage mean that a large proportion of patients with OP have never undergone DXA examination.<sup>9</sup> An alternative method, quantitative computed tomography (qCT), also faces challenges due to its high operational cost and limited feasibility for large-scale screening.<sup>61</sup>

Over the past few decades, the rapid rise of AI technologies has provided a promising solution to these diagnostic challenges. With powerful image recognition capabilities, AI systems offer high accuracy, rapid processing speed, and cost-effectiveness, making it feasible to diagnose OP using routine imaging modalities.<sup>62</sup> Beyond imaging-based diagnosis, risk factor screening and prediction models powered by AI have also become active areas of current research, offering complementary tools for early and noninvasive identification of OP.

## Image Recognition

AI possesses powerful image recognition capabilities and has been applied across multiple imaging modalities.<sup>63</sup> In the field of OP, numerous studies have explored AI-assisted diagnostic approaches based on medical image analysis.

### X-Ray Imaging

Different imaging modalities contribute unequally to AI-based OP diagnosis. X-ray imaging has emerged as the most extensively studied modality due to its high accessibility, low cost, and widespread clinical use.<sup>64</sup> However, unlike dual-energy DXA, conventional X-rays are not sufficiently sensitive to detect OP, limiting their diagnostic utility. With advances in AI, especially in ML and DL, it has become possible to extract subtle image features imperceptible to human observers. That enabling the construction of large-scale datasets required for robust AI training.<sup>65</sup> Consequently, a larger body of literature has focused on X-ray–based approaches, whereas studies involving MRI and US remain comparatively limited. The use of standard X-rays for OP diagnosis has gained increasing attention, particularly over the past five years (Table 1). Among studies applying AI to X-ray images for OP diagnosis, the most commonly used anatomical sites include the chest,<sup>66</sup> spine,<sup>67</sup> pelvis and hip,<sup>68</sup> hands<sup>69</sup> and feet,<sup>70</sup> and or maxillofacial region.<sup>47</sup>

Chest radiographs are among the most common medical images, and several studies have investigated their utility in OP diagnosis. For example, Lin et al and Asamoto et al both developed DL models trained on chest X-rays to detect OP.

**Table I** Recent Studies Applying AI for OP Diagnosis Using X-Ray Imaging

| Parts | Researchers                      | Year | Purpose                                                                                                                                              | Algorithms Evaluated                                        | Dataset                               | Best-Performing Algorithm | Performance Metrics of Best-Performing Algorithm |
|-------|----------------------------------|------|------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------|---------------------------------------|---------------------------|--------------------------------------------------|
| Knee  | Naguib SM et al <sup>71</sup>    | 2024 | Develop a new deep-learning approach to categorize knee X-ray images into OP, osteopenia, and normal classes                                         | AlexNet, ResNet50                                           | Knee X-ray images                     | Superfluity DL model      | AUC0.8116(CI: Not given)                         |
|       | Xie H et al <sup>72</sup>        | 2024 | Develop a FSL framework for the diagnosis of osteopenia and OP in knee X-ray images                                                                  | VGG, ResNet, Xception                                       | Knee X-ray images, chest x-ray images | FSL models                | Accuracy0.728(CI: Not given)                     |
|       | Sarmadi A et al <sup>73</sup>    | 2024 | Compare the ability of OP detection between vision transformer and neural networks                                                                   | CNN, ViT                                                    | Knee X-ray images                     | vit_r50_l32_fe            | Accuracy0.6380(CI: Not given)                    |
|       | Srinivasan D et al <sup>74</sup> | 2025 | Improve the accuracy, speed, and reliability of early-stage OP detection by ANNs                                                                     | ANN                                                         | Knee X-ray images                     | BoneVoyage model          | Accuracy0.972(CI: Not given)                     |
|       | Muhammad I et al <sup>75</sup>   | 2025 | Introduce an advanced deep-learning methodology to enhance the accuracy and efficiency of osteoporosis detection through knee X-ray analysis         | DenseNet169, Vision Transformer (ViT), Attention Model (AM) | Knee X-ray images                     | Fusion model              | Accuracy0.8611(CI: Not given)                    |
|       | Qureshi MB et al <sup>76</sup>   | 2025 | Detect OP from knee X-rays using DL, specifically employing the ResNet-50 model with transfer learning, as a strategy to address existing challenges | VGG-16, ResNet-18, ResNet-50                                | Knee X-ray images                     | ResNet-50                 | Accuracy0.90(CI: Not given)                      |

|       |                               |      |                                                                                                                                       |                                 |                    |             |                                                                                                                        |
|-------|-------------------------------|------|---------------------------------------------------------------------------------------------------------------------------------------|---------------------------------|--------------------|-------------|------------------------------------------------------------------------------------------------------------------------|
| Chest | Jang M et al <sup>66</sup>    | 2021 | Develop and evaluate the DL approaches for screening OP using the classified chest radiographs based on the DXA-based diagnosis of OP | CNN                             | Chest X-ray images | OsPor       | Internal validation AUC 0.91 (95%CI: 0.90–0.92), External validation AUC 0.88 (95%CI: 0.85–0.90)                       |
|       | Lin C et al <sup>77</sup>     | 2024 | Evaluate the effectiveness of DXA screening in high-risk patients with OP identified by an AI model using chest radiographs.          | CNN                             | Chest X-ray images | DenseNet    | Positive predictive value 75% (CI: Not given)                                                                          |
|       | Asamoto T et al <sup>78</sup> | 2024 | Verify the usefulness of DL model for predicting BMD of femoral neck in external validation                                           | ResNet-50                       | Chest X-ray images | ResNet-50   | Osteopenia AUC 0.80, OP AUC 0.83 (CI: Not given)                                                                       |
|       | Tsai DJ et al <sup>79</sup>   | 2024 | Develop a DL model to identify OP from chest X-ray features                                                                           | DenseNet                        | Chest X-ray images | CXR-OP      | Internal validation AUC 0.930 (CI: Not given), External validation AUC 0.892 (CI: Not given)                           |
|       | Ohta Y et al <sup>80</sup>    | 2025 | Develop and evaluate two DL models to detect low BMD in the femoral neck and lumbar vertebrae                                         | ResNet-50                       | Chest X-ray images | ResNet-50   | Femoral neck AUC 0.82 (CI: Not given); Lumbar vertebrae AUC 0.96 (CI: Not given)                                       |
|       | Tang J et al <sup>81</sup>    | 2025 | Provide an automatic screening method for OP using chest X-rays                                                                       | Inception v3, VGG-16, ResNet-50 | Chest X-ray images | ResNet-50   | Internal validation AUC 0.945 (CI: Not given) and 0.957 (CI: Not given), External validation AUC 0.904 (CI: Not given) |
|       | Iwao Y et al <sup>82</sup>    | 2025 | Predicts BMD from chest radiographs using clavicular features                                                                         | U-Net                           | Chest X-ray images | U-Net       | Accuracy 0.68 (CI: Not given)                                                                                          |
|       | Park J et al <sup>83</sup>    | 2025 | Develop and validate a DL model for the opportunistic detection of low bone mass using routine chest X-rays                           | OsPenScreen                     | Chest X-Ray Images | OsPenScreen | AUC 0.95 (95% CI: 0.94–0.97)                                                                                           |

(Continued)

Table I (Continued).

| Parts | Researchers                  | Year | Purpose                                                                                                                                                                          | Algorithms Evaluated                                                     | Dataset                                | Best-Performing Algorithm                              | Performance Metrics of Best-Performing Algorithm                                                                                                                                                                   |
|-------|------------------------------|------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------|----------------------------------------|--------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Spine | Zhang B et al <sup>67</sup>  | 2020 | Develop a DCNN model to classify osteopenia and OP with the use of lumbar spine X-ray images                                                                                     | DCNN                                                                     | Lumbar spine X-ray images              | DCNN                                                   | Test dataset 1 OP AUC 0.767(95%CI: 0.701–0.824), Test dataset 1 osteopenia AUC 0.787(95%CI: 0.701–0.824); Test dataset 2 OP AUC 0.726(95%CI: 0.646–0.796), Test dataset 2 osteopenia AUC 0.810(95%CI: 0.737–0.870) |
|       | Lee S et al <sup>84</sup>    | 2020 | Evaluated ML models for identifying individuals with abnormal BMD through an analysis of spine X-ray features                                                                    | AlexNet, VGG, Inception-V3, ResNet-50, SVMC, KNNC, RFC                   | Spine X-ray images                     | RFC with VGG-I6                                        | AUC0.74(CI: Not given)                                                                                                                                                                                             |
|       | Hsieh CI et al <sup>85</sup> | 2021 | Present an automated tool to identify fractures, predict BMD, and evaluate fracture risk using plain radiographs                                                                 | VGG-11, VGG-16, ResNet-18, ResNet-34                                     | Pelvis/ lumbar spine X-ray images      | VGG-16, ResNet-34                                      | Hip OP AUC 0.89(CI: Not given);Spine OP AUC 0.89(CI: Not given)                                                                                                                                                    |
|       | Mao L et al <sup>86</sup>    | 2022 | Construct a CNN model for screening primary osteopenia and OP based on the lumbar radiographs                                                                                    | DenseNet                                                                 | Lumbar X-ray images                    | The model based on the anteroposterior\lateral channel | AUC0.909(95%CI: 0.909–0.950),0.937(95% CI: 0.883–0.930)                                                                                                                                                            |
|       | Dong Q et al <sup>87</sup>   | 2023 | Develop a DL classifier for spinal osteoporotic compression fractures                                                                                                            | GoogLeNet                                                                | Spine X-Ray images                     | GoogLeNet                                              | AUC0.99(95%CI: 0.98–1.00)                                                                                                                                                                                          |
|       | Hong N et al <sup>88</sup>   | 2023 | Develop DL scores to detect OP and VF based on lateral spine radiography and investigate whether their use can improve referral of high-risk individuals to bone-density testing | EfficientNet-B4                                                          | Spine X-Ray images                     | VERTE-X pVF, VERTE-X                                   | Vertebral fractures AUC 0.926(95%CI: 0.908–0.944); OP AUC 0.848(95%CI: 0.827–0.869)                                                                                                                                |
|       | Dong Q et al <sup>89</sup>   | 2024 | Build an automated opportunistic OCF radiograph screening tool with three primary sequential components                                                                          | GoogLeNet, Inception-ResNet-v2, EfficientNet-B1, two ensemble algorithms | Thoracic and lumbar spine X-ray images | Ensemble Model                                         | Local dataset's AUC 0.948(95%CI: 0.933–0.961),MrOS dataset's AUC 0.936(95%CI: 0.910–0.959)                                                                                                                         |
|       | Moro T et al <sup>90</sup>   | 2025 | Develop an AI-assisted diagnostic system that estimated not only lumbar BMD but also femoral BMD from anteroposterior lumbar X-ray images                                        | ANN                                                                      | Lumbar X-ray images                    | ANN                                                    | AUC>0.8(CI: Not given)                                                                                                                                                                                             |

|       |                                  |      |                                                                                                                                                                                   |                                                      |                            |                                                                                                                                       |                                                                                                 |
|-------|----------------------------------|------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------|----------------------------|---------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------|
|       | Hong N et al <sup>91</sup>       | 2025 | Develop DL models to detect prevalent vertebral fracture and OP                                                                                                                   | CNN                                                  | Lateral spine X-Ray images | VERTE-X                                                                                                                               | VERTE-X OP AUC 0.926 (95% CI 0.908–0.955), 0.848 (95% CI 0.827–0.869)                           |
|       | Hong J et al <sup>92</sup>       | 2025 | Develop a DL -based model to predict pediatric BMD using plain spine radiographs                                                                                                  | ResNet-18                                            | Spine X-Ray images         | ResNet-18                                                                                                                             | AUC0.85(CI: Not given)                                                                          |
|       | Tamai K et al <sup>93</sup>      | 2025 | Validate the diagnostic yield of a DL algorithm for detecting osteopenia/ OP using cervical radiography and compare its diagnostic accuracy with that of spine surgeons           | EfficientNetB2                                       | Cervical spine radiography | EfficientNetB2                                                                                                                        | AUC0.869 (95% CI: 0.822–0.916)                                                                  |
|       | Boonrod A et al <sup>94</sup>    | 2025 | Propose the DL models to screen for OP and measure its accuracy on OP detection from lumbar spine radiographs                                                                     | MobileNet-V2, AlexNet, DarkNet-19, ResNet-18, CiRA32 | Lumbar X-ray images        | ResNet-18                                                                                                                             | AUC0.79 (95%CI 0.76–0.82)                                                                       |
| Mouth | Kavitha MS et al <sup>47</sup>   | 2015 | Determine whether individual measurements or a combination of textural features and mandibular cortical width (MCW) derived from digital DPRs are more useful in assessment of OP | k-NN, SVM, NB                                        | Panoramic radiography      | SVM                                                                                                                                   | Femoral neck BMD accuracy 0.963(CI: Not given); Lumbar spine BMD accuracy 0.958 (CI: Not given) |
|       | Kavitha MS et al <sup>95</sup>   | 2016 | Propose a new automated screening system based on a hybrid genetic swarm fuzzy (GSF) classifier using digital DPRs to diagnose females with a low BMD or OP                       | GSF classifier                                       | Panoramic radiographs      | Hybrid GSF with optimized MF and RS                                                                                                   | Lumbar spine accuracy 0.9601 (95% CI: 0.903–0.995) Femoral neck accuracy 0.989 (95% CI:0.920–1) |
|       | Hwang JJ et al <sup>96</sup>     | 2017 | Develop an OP detection model based on panoramic radiography                                                                                                                      | decision tree, SVM                                   | Panoramic radiography      | SVM                                                                                                                                   | Accuracy 0.969(CI: Not given)                                                                   |
|       | Lee JS et al <sup>97</sup>       | 2018 | Evaluate the diagnostic performance of a DCNN-based computer-assisted diagnosis (CAD) system in the detection of OP on panoramic radiographs                                      | DCNN                                                 | Panoramic radiographs      | SC-DCNN                                                                                                                               | AUC0.9991(CI: Not given)                                                                        |
|       | Alzubaidi MA et al <sup>98</sup> | 2019 | Explores the use of 13 different feature extractors for detection of reduced bone density in dental radiographs                                                                   | SVM                                                  | Panoramic radiographs      | The models based on SOM and LVQ using Gabor Filter, Edge Orientation Histogram, Haar Wavelet, and Steerable Filter feature extractors | Accuracy 0.926(CI: Not given)                                                                   |

(Continued)

Table I (Continued).

| Parts | Researchers                     | Year | Purpose                                                                                                                  | Algorithms Evaluated                                  | Dataset               | Best-Performing Algorithm         | Performance Metrics of Best-Performing Algorithm                                                                                                                                                         |
|-------|---------------------------------|------|--------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------|-----------------------|-----------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|       | Nakamoto T et al <sup>99</sup>  | 2022 | Develop computer-aided screening systems that could predict OP                                                           | AlexNet, VGG-16, GoogLeNet                            | Panoramic radiographs | GoogLeNet                         | Lumbar spine OP accuracy 0.74(95% CI:0.654–0.823),0.79(95% CI:0.710–0.870),0.79(95% CI:0.710–0.870). Femoral neck OP accuracy 0.70(95% CI:0.610–0.790),0.75(95% CI:0.665–0.835),0.75(95% CI:0.665–0.835) |
|       | Tassoker M et al <sup>100</sup> | 2022 | Compare five CNN for predicting OP based on mandibular cortical index (MCI) on panoramic radiographs                     | AlexNet, GoogleNet, ResNet-50, SqueezeNet, ShuffleNet | Panoramic radiographs | AlexNet                           | C1, C3 AUC 0.9987±(0.0012) (CI: Not given)                                                                                                                                                               |
|       | Sukegawa S et al <sup>101</sup> | 2022 | Investigate the use of DL to classify OP from DPRs                                                                       | ResNet, EfficientNet                                  | Panoramic radiographs | Ensemble model of EfficientNet-b7 | AUC0.921 (95%CI:0.917–0.925)                                                                                                                                                                             |
|       | Yadalam PK et al <sup>102</sup> | 2025 | Classify bone loss severity (normal, mild, or severe) from OPG notes using a novel dual-embedding FSL framework          | DualFit                                               | Panoramic radiographs | DualFit                           | Accuracy of 0.9898(CI: Not given)                                                                                                                                                                        |
|       | Roebbers P et al <sup>103</sup> | 2025 | Investigate the use of NN to predict potential OP using the Panoramic Mandibular Index derived from panoramic radiograph | ResNet-50                                             | Panoramic radiographs | ResNet-50                         | Accuracy 0.895(CI: Not given)                                                                                                                                                                            |

|                |                                       |      |                                                                                                                                                                                          |                                                                   |                             |                          |                                                                              |
|----------------|---------------------------------------|------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------|-----------------------------|--------------------------|------------------------------------------------------------------------------|
| Hip            | Sapthagirivasan V et al <sup>68</sup> | 2013 | Evaluate the ability of a kernel-based SVM with respect to diagnosis and add to knowledge about the trabecular features of digital hip radiographs for identifying subjects with low BMD | SVM                                                               | Hip X-ray images            | SVM                      | Accuracy 0.90 (95%CI:0.82–0.98)                                              |
|                | Yamamoto N et al <sup>104</sup>       | 2020 | Consider the use of DL to diagnose OP from hip radiographs, and whether adding clinical data improves diagnostic performance over the image mode alone                                   | ResNe-t18, ResNet-34, GoogleNet, EfficientNet b3, EfficientNet b4 | Hip X-ray images            | EfficientNet b3          | AUC 0.9374(CI: Not given)                                                    |
|                | Jang R et al <sup>105</sup>           | 2021 | Predict OP via simple hip radiography using DL algorithm                                                                                                                                 | VGG-16                                                            | Hip X-ray images            | VGG-16                   | AUROC0.867(CI: Not given)                                                    |
|                | Yamamoto N et al <sup>106</sup>       | 2021 | Evaluate the OP identification ability of DL by combining hip radiographs with patient variables                                                                                         | ResNet-18, ResNet-34, ResNet-50, ResNet-101, ResNet-152           | Hip X-ray images            | ResNet-34 ensemble model | AUC0.892 (95% CI: 0.890–0.894)                                               |
|                | Nguyen TP et al <sup>107</sup>        | 2021 | Proposing a DL -based method for evaluation of BMD based on the X-rays of hips and biological parameters.                                                                                | CNN                                                               | Hip X-ray images            | CNN                      | Correlation coefficient 0.8075(CI: Not given)                                |
|                | Acosta-Batlle J et al <sup>108</sup>  | 2025 | Develop and validate an AI-based tool for the diagnosis of OP                                                                                                                            | CNN                                                               | Hip X-ray images            | CNN                      | AUC0.866(CI: Not given)                                                      |
| Hands and feet | Areeckal AS et al <sup>69</sup>       | 2017 | Propose an automated low-cost tool for early diagnosis of onset of OP using cortical radiogrammetry and cancellous texture analysis from hand and wrist radiographs                      | ANN                                                               | Hand and wrist X-ray image  | ANN                      | Training accuracy 0.943 (CI: Not given); Test accuracy 0.885 (CI: Not given) |
|                | Singh A et al <sup>109</sup>          | 2017 | Extract discriminatory statistical features of different orders used for ML for the classification of two populations composed of osteoporotic and healthy subjects                      | SVM, NB, ANN, k-NN                                                | Calcaneus X-ray image       | SVM                      | Classification rate 98% (CI: Not given)                                      |
|                | Gharehmohammadi F et al <sup>70</sup> | 2025 | Create a DL model to screen for OP /osteopenia in patients by using radiographs of the foot or ankle                                                                                     | CNN, FCNN                                                         | X-rays of the foot or ankle | CNN                      | AUC 0.87(95% CI:0.85–0.94)                                                   |

(Continued)

Table 1 (Continued).

| Parts  | Researchers                      | Year | Purpose                                                                                                                                      | Algorithms Evaluated                | Dataset                           | Best-Performing Algorithm | Performance Metrics of Best-Performing Algorithm                                                |
|--------|----------------------------------|------|----------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------|-----------------------------------|---------------------------|-------------------------------------------------------------------------------------------------|
|        | Ho CS et al <sup>57</sup>        | 2025 | Develop a HarDNet-based DL regression model for hand BMD prediction.                                                                         | CNN                                 | Hand X-ray images                 | DeepDXA-Hand              | Osteopenia accuracy 0.82 (95% CI:0.80–0.84); OP accuracy 0.80(95% CI:0.78, 0.82)                |
|        | Hwang U et al <sup>110</sup>     | 2025 | Propose a method to predict OP using hand and wrist X-ray images                                                                             | U-Net, SimCLR, SupCon, SwAV, VICReg | Hand X-ray images                 | SimCLR                    | AUC0.81(CI: 0.78–0.84)                                                                          |
|        | Lee KA et al <sup>111</sup>      | 2026 | Perform segmentation and classification on metacarpal radiographs to accurately detect OP                                                    | OMO-Net                             | Hand X-ray images                 | OMO-Net                   | AUC0.99(CI: Not given)                                                                          |
| Others | Liu J et al <sup>112</sup>       | 2019 | Present an improved OP diagnosis algorithm based on U-NET network                                                                            | U-Net                               | Pelvic X-ray image                | U-Net                     | SO-S1 AUC 0.6921(CI: Not given);SO-S2 AUC 0.7011(CI: Not given);S1-S2 AUC 0.6348(CI: Not given) |
|        | Zhou K et al <sup>113</sup>      | 2025 | Explore the feasibility of using a hybrid DL framework (HDLF) to establish a model for BMD prediction and classification based on BPX images | CNN                                 | Biplanar X-ray radiography        | CNN                       | AUC0.97(CI: Not given)                                                                          |
|        | Yen TY et al <sup>114</sup>      | 2024 | Explore the possibility of predicting the BMD and classifying high-risk patient groups using KUB radiographs                                 | DeepDXA-KUB (a DL model)            | Kidney-ureter-bladder radiographs | DeepDXA-KUB(a DL model)   | Lumbar vertebrae AUC 0.939(CI: Not given); Left hip AUC 0.947(CI: Not given)                    |
|        | Quagliato L et al <sup>115</sup> | 2025 | Develop an ML algorithm that performs binary classification of OF risk based on 2D-DXA-derived geometry and material parameters              | XGBoost, SVM, MLP                   | 2D-DXA                            | XGBoost                   | Raw data AUC0.77(CI: Not given), Synthetic data AUC 0.99(CI: Not given)                         |

**Abbreviations:** AI, artificial intelligence; AM, Attention Model; ANN, artificial neural network; BMD, bone mineral density; BPX, biplanar X-ray imaging; CAD, computer-aided diagnosis; CI, confidence interval; CNN, convolutional neural network; DCNN, deep convolutional neural network; DL, deep learning; DXA, dual-energy X-ray absorptiometry; FSL, few-shot learning; GSF, genetic swarm fuzzy; HDLF, hybrid deep learning framework; HU, Hounsfield unit; k-NN, k-nearest neighbors; KNNC, k-nearest neighbors classifier; LR, logistic regression; LVQ, learning vector quantization; MCW, mandibular cortical width; MF, membership function; ML, machine learning; MLP, multilayer perceptron; NB, naïve Bayes; OCF, osteoporotic compression fracture; OF, osteoporotic fracture; OP, osteoporosis; OPG, orthopantomogram; OVf, osteoporotic vertebral fracture; ResNet, residual neural network; RFC, random forest classifier; ROI, region of interest; RS, rule set; SC-DCNN, single-channel deep convolutional neural network; SOM, self-organizing map; SVM, support vector machine; SVMC, support vector machine classifier; U-Net, U-shaped convolutional neural network; ViT, vision transformer; VF, vertebral fracture; AUC, area under the receiver operating characteristic curve; AUROC, area under the receiver operating characteristic curve; ACC, accuracy.

Lin et al employed a CNN based on the DenseNet architecture, while Asamoto et al integrated imaging data with patient age and sex. Both models achieved diagnostic accuracies above 75%, and Asamoto's group further externally validated a femoral BMD prediction model with similar accuracy approaching 80%.<sup>77,78</sup> Jang et al developed a supervised DL model named OsPor-Screen, which demonstrated even higher performance, achieving AUCs of 0.91 and 0.88 on internal and external test datasets, respectively highlighting the strong potential of chest X-rays for OP screening.<sup>66</sup>

Dental panoramic radiographs (DPRs) have also been widely explored for OP detection, with research in this area dating back nearly a decade. In 2015, Kavitha et al used DPRs to train NB, k-Nearest Neighbors (k-NN), and SVM models, utilizing fractal dimension (FD) and gray-level co-occurrence matrix (GLCM) features to classify OP. All three models achieved accuracies exceeding 90%,<sup>47</sup> and subsequent model refinements increased performance to over 95%.<sup>95</sup> With the advent of DL, Lee et al (2018) trained Deep Convolutional Neural Network (DCNN) models with two architectures: a single-column DCNN (SC-DCNN) using a region of interest (ROI) below the mandible, and a multi-column DCNN (MC-DCNN) using bilateral mandibular ROIs. The SC-DCNN achieved an accuracy of 92.5%, SC-DCNN (with data augmentation) 98%, and MC-DCNN 98.5%.<sup>97</sup> Subsequent studies by Sukegawa,<sup>101</sup> Tassoker,<sup>100</sup> and Nakamoto<sup>99</sup> applied pretrained models such as AlexNet and GoogLeNet. Notably, Nakamoto et al extended their trained models to lumbar spine and femoral neck X-rays, achieving diagnostic performance comparable to that of experienced radiologists. A distinct line of research has focused on feature extraction from DPRs.<sup>99</sup> For instance, Alzubaidi et al utilized 13 feature extractors, including pixel intensity histograms, and trained Self-Organizing Map (SOM) and Learning Vector Quantization (LVQ) models. Among them, SOM/LVQ models incorporating Gabor filters, edge orientation histograms, Haar wavelets, and steerable filters demonstrated the best OP detection accuracy.<sup>98</sup>

Spinal radiographs are widely used in orthopedic diagnostics and have become a popular research focus in AI-based OP detection. Many studies have employed pretrained CNN architectures, such as those by Lee et al<sup>84</sup> and Dong et al<sup>87</sup> demonstrating the feasibility of spinal image-based OP recognition. Other studies have applied deep DCNNs to spine imaging, including works by Zhang et al<sup>67</sup> and Hong et al<sup>88</sup>. Notably, Hong's team explored whether integrating clinical parameters (such as age, sex, or biochemical markers) with DCNN-derived imaging features could improve diagnostic performance. Their findings confirmed that the combination of imaging and clinical data yielded superior diagnostic accuracy for OP compared with imaging alone.<sup>88</sup> Recent research has advanced beyond OP diagnosis alone, using DL models to simultaneously identify OP, vertebral fractures, and fracture risk from spinal radiographs.<sup>91</sup> These developments suggest that DL-based spinal imaging analysis may eventually expand to the diagnosis of other spinal disorders as well.

Pelvic and hip radiographs have also become major areas of interest for AI-based OP diagnosis. AI analysis of hip X-rays began relatively early: in 2013, Sapthagirivasan et al applied an SVM-based model that extracted trabecular bone features from hip radiographs, achieving an average accuracy of 90%.<sup>68</sup> Later, DL methods were introduced into this field. Liu et al employed a U-Net model, though its accuracy was limited,<sup>112</sup> while Yamamoto et al achieved improved results using pretrained CNNs, surpassing earlier traditional ML performance.<sup>104</sup> Building on these efforts, Nguyen et al integrated a Sobel gradient-based CNN mapping model with biological parameters to predict OP, achieving a correlation coefficient of 0.8075 compared with DXA measurements, indicating strong predictive capability.<sup>107</sup> Furthermore, Srinivasan et al developed a dual-core model named BoneVoyage for hip radiographs, which combines ShuffleNet for efficient feature extraction with artificial neural networks (ANNs) for classification. This hybrid model achieved an impressive 97.2% accuracy, significantly outperforming conventional diagnostic approaches and pointing toward a promising direction for future multimodal AI models.<sup>74</sup>

Research on knee X-rays for OP diagnosis has been relatively limited compared with other imaging modalities, largely because such images are more commonly used for evaluating osteoarthritis rather than bone density. However, studies published in 2024 by Sarmadi et al,<sup>73</sup> Xie et al,<sup>72</sup> and Naguib et al<sup>71</sup> have renewed interest in this area. These researchers primarily employed pretrained DL models, with notable innovation by Xie et al, who implemented a few-shot learning (FSL) approach to address the challenge of limited imaging data. Their results demonstrated that the FSL model achieved higher accuracy and sensitivity than radiologists, underscoring the potential of AI even in small-sample OP datasets.<sup>72</sup> This also provides a potential approach to addressing the algorithmic fairness issues that inherently exist in the use of artificial intelligence for disease diagnosis.<sup>116</sup> Compared with knee images, hand and foot X-rays are more complex due to the intricate bone structures and overlapping features, making feature extraction challenging.

Nonetheless, advances in AI have made it possible to utilize these images for OP diagnosis. In 2017, Singh et al applied four ML algorithms—SVM, k-NN, NB, and ANNs—to analyze trabecular bone features in calcaneal X-rays. All classifiers achieved accuracies above 95%, with SVM performing best (97.87% accuracy).<sup>109</sup> More recently, Ho et al (2025) developed an advanced DL model named DeepDXA-Hand, which analyzed multiple hand regions—including the capitate, trapezoid, hamate, triquetrum, and second metacarpal head—and achieved an overall accuracy of 0.80.<sup>57</sup> This model demonstrated the feasibility of using AI to perform noninvasive OP screening based on standard hand radiographs.

Beyond the imaging sites above, several studies have explored the potential of alternative X-ray modalities for OP detection. For instance, Zhao et al used ML to diagnose OP from shoulder radiographs;<sup>117</sup> Yen et al developed a CNN-based model for detecting OP from kidney–ureter–bladder (KUB) X-rays;<sup>114</sup> Zhou et al applied DL to biplanar radiography for bone density estimation;<sup>113</sup> and Ashok Kumar et al utilized forearm X-rays to predict the risk of future OF in women.<sup>118</sup> Although the number of studies in these areas remains limited, they collectively illustrate the growing versatility and potential of AI in diverse radiographic applications for OP diagnosis.

In summary, due to its high accessibility, low cost, and widespread clinical use, X-ray imaging,<sup>64</sup> when combined with SVM or pretrained DL models (such as VGG and ResNet) for feature extraction and classification, enables a low-cost, simple, and automated diagnosis of OP patients.<sup>85,95</sup>

## CT

CT is a widely used imaging modality in clinical diagnosis, known for its high spatial resolution and three-dimensional (3D) imaging capability. A notable derivative of CT technology, qCT, has been recognized as a potential alternative to DXA for diagnosing OP.<sup>119</sup> Current AI-based CT studies on OP mainly focus on chest,<sup>120</sup> abdominal,<sup>121</sup> spinal,<sup>122</sup> and oromaxillofacial<sup>123</sup> regions, with an increasing number of investigations also aiming to enhance the diagnostic performance of qCT<sup>124</sup> (Table 2).

Although chest, abdominal, and spinal CT scans are frequently performed in routine clinical practice, they are not traditionally used for OP diagnosis, because they cannot directly visualize reductions in BMD. However, with the advent of AI, it has become feasible to detect OP using these common CT datasets.<sup>168</sup> Most AI approaches follow a two-step workflow: (1) vertebral segmentation and (2) feature extraction and classification.<sup>63, 169</sup>

For vertebral segmentation, various methods have been proposed. Asaka et al manually extracted L1–L4 vertebral levels from unenhanced abdominal CT scans to train a CNN model, achieving an AUC of 0.965 (internal) and 0.970 (external validation), demonstrating excellent diagnostic accuracy.<sup>122</sup> Similar frameworks were employed by Tariq et al<sup>140</sup> and Tomita et al<sup>120</sup> More recent studies have adopted automatic segmentation using DL architectures. For instance, Fang et al used U-Net for automatic segmentation followed by DenseNet-121 for BMD estimation, achieving a very high correlation with qCT-measured BMD ( $r > 0.98$ ).<sup>158</sup> Comparable approaches have been reported by Oh,<sup>41</sup> Pan,<sup>155</sup> and Li,<sup>56</sup> all employing U-Net-based pipelines. Alternative models have also been explored: Breit et al proposed a CNN using wavelet-based and geometric constraints for segmentation;<sup>126</sup> Pan et al (2023) utilized a landmark detection network combining Single Shot Multibox Detector (SSD) and VGG-16;<sup>156</sup> Peng et al used a VB-Net–based pretrained model;<sup>127</sup> and other studies have applied Vision Transformers.<sup>136</sup> Although 2D segmentation is possible, Hathaway et al demonstrated that 3D segmentation provides superior accuracy.<sup>157</sup>

Following segmentation, CNNs are commonly employed for OP classification. Dzierżak et al compared six transfer learning models—VGG16, VGG19, MobileNetV2, Xception, ResNet50, and InceptionResNetV2—to address small-sample challenges, with VGG16 performing best (accuracy 95%).<sup>146</sup> Tang et al developed an automated CNN framework for lumbar CT analysis, comprising a Mark-Segmentation Network (MS-Net) and a BMD-Classification Network (BMDC-Net). The model achieved a classification accuracy of 76.65% on the test dataset, demonstrating strong potential for automated OP diagnosis.<sup>145</sup>

With the rise of radiomics, this technique has become increasingly relevant in CT-based OP research. A typical radiomics workflow involves five steps:

- (1) Image Acquisition: Single-source dual-energy CT scans generate 70-keV virtual monochromatic images of the lumbar spine with a bone-density calibration phantom.

**Table 2** Applications of AI in OP-Related Tasks Based on CT Imaging

| Task                   | Parts | Researchers                   | Year | Purpose                                                                                                                                                                                                                                | Algorithms Evaluated                                                                                                                                                                                                                                                                                                                                                                                      | Dataset                         | Performance Metrics                                                                                                                                                                                                        |
|------------------------|-------|-------------------------------|------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Osteoporosis Diagnosis | Chest | Yang J et al <sup>125</sup>   | 2022 | Explore the value of the attenuation values of all thoracic vertebrae and the first lumbar vertebra measured by AI on non-enhanced chest CT to do OP screening                                                                         | AI-Rad Companion                                                                                                                                                                                                                                                                                                                                                                                          | Chest CT                        | Osteopenia AUC0.831(95% CI: 0.800–0.862); OP AUC0.972(95% CI: 0.961–0.983)                                                                                                                                                 |
|                        | Chest | Breit HC et al <sup>126</sup> | 2023 | Assess the performance of an AI-based CNN-Algorithm for the detection of low bone density on routine non-contrast chest CT                                                                                                             | CNN                                                                                                                                                                                                                                                                                                                                                                                                       | Chest CT                        | Accuracy0.75(CI: Not given)                                                                                                                                                                                                |
|                        | Chest | Peng T et al <sup>127</sup>   | 2023 | Explore the feasibility of using DL to establish a model for OP classification and bone density value prediction based on opportunistic CT scans and to verify its generalization and diagnostic ability using an independent test set | DenseNet, VB-Net                                                                                                                                                                                                                                                                                                                                                                                          | Opportunistic CT                | Normal AUC0.999(95% CI: 0.999–1.000); Osteopenia AUC0.970(95% CI: 0.949–0.990); OP AUC0.933(95% CI: 0.906–0.960)                                                                                                           |
|                        | Chest | Li J et al <sup>56</sup>      | 2024 | Develop a DL model to automatically measure BMD and improve the diagnostic rate of OP                                                                                                                                                  | GCN, 3D-Unet, 3D-ResNet                                                                                                                                                                                                                                                                                                                                                                                   | Chest or abdominal paired CT    | Accuracy0.88(CI: Not given); 0.91(CI: Not given); 0.91(CI: Not given)                                                                                                                                                      |
|                        | Chest | Park H et al <sup>128</sup>   | 2024 | Examine the diagnostic efficacy of automated deep learning-based bone mineral density (DL-BMD) measurements for OP                                                                                                                     | 2D U-Net                                                                                                                                                                                                                                                                                                                                                                                                  | Chest CT, abdomen CT, spine CT  | Low BMD AUC0.790(95% CI 0.733–0.839); OP AUC0.769(95% CI 0.710–0.820)                                                                                                                                                      |
|                        | Chest | Pan J et al <sup>58</sup>     | 2024 | Develop a multi-feature DL model based on chest CT, combined with clinical information and radiomics to explore the feasibility in screening for OP based on estimation of volumetric BMD                                              | DCNN                                                                                                                                                                                                                                                                                                                                                                                                      | Chest CT                        | Normal AUC0.992(CI: Not given), Osteopenia AUC0.973(CI: Not given), OP AUC0.989(CI: Not given)                                                                                                                             |
|                        | Chest | Lin X et al <sup>129</sup>    | 2024 | Investigate the feasibility of utilizing radiomics technology based on chest CT images to screen for opportunistic OP                                                                                                                  | SVM                                                                                                                                                                                                                                                                                                                                                                                                       | Chest CT                        | Internal AUC0.892(95% CI 0.8399–0.9445); External AUC0.7960(95% CI 0.7361–0.8563)                                                                                                                                          |
|                        | Chest | Fang K et al <sup>130</sup>   | 2024 | Assess the possibility of using radiomics, DL, and transfer learning methods for the analysis of chest CT scans and combine these techniques with bone turnover markers to identify and screen for OP                                  | SVM, KNN, RandomForest, ExtraTrees, XGBoost, LightGBM, NaiveBayes, AdaBoost, GradientBoosting, LR, MLP, alexnet, densenet121, densenet169, googlenet, mnasnet1_0, mobilenet_v2, mobilenet_v3_large, mobilenet_v3_small, resnet101, resnet152, resnet18, resnet34, resnet50, resnext50_32x4d, squeezeNet1_0, squeezeNet1_1, vgg11, vgg11_bn, vgg13, vgg13_bn, vgg16, vgg16_bn, vgg19, vgg19_bn, shuffleNet | Chest CT                        | Best performance:3D DL model; AUC0.906(95% CI 0.846–0.966)                                                                                                                                                                 |
|                        | Chest | Tong X et al <sup>131</sup>   | 2024 | Develop two bone status prediction models combining DL and radiomics based on standard-dose chest computed tomography (SDCT) and LDCT                                                                                                  | VB-Net, RF                                                                                                                                                                                                                                                                                                                                                                                                | SDCT, LDCT                      | LDCT AUC0.97 ± 0.01(CI: Not given);SDCT AUC0.94 ± 0.02(CI: Not given)                                                                                                                                                      |
|                        | Chest | Wang S et al <sup>132</sup>   | 2024 | Develop an intelligent diagnostic model for OP screening based on LDCT                                                                                                                                                                 | SVM, RF                                                                                                                                                                                                                                                                                                                                                                                                   | LDCT                            | Best performance: Aseg model; First-level model: internal AUC0.985(CI: Not given); external internal AUC0.965(CI: Not given).In the second-level model: internal AUC0.933(CI: Not given); external AUC0.907(CI: Not given) |
|                        | Chest | Li Y et al <sup>133</sup>     | 2025 | Explore the feasibility of DL -enhanced, fully automated BMD measurement using the ultralow-voltage 80 kV chest CT scans                                                                                                               | DenseNet, ResNet                                                                                                                                                                                                                                                                                                                                                                                          | 80kV chest CT, 120kV lumbar CT  | Best performance:80 kV CNNs; OP AUC0.997–1.000 (CI: Not given);Low BMD AUC0.997–0.998(CI: Not given)                                                                                                                       |
|                        | Chest | Li Y et al <sup>134</sup>     | 2025 | Propose a DL method for osteoporosis diagnose using LDCT and lumbar CT scans across CT scanners from different manufacturers and hospitals                                                                                             | CNN                                                                                                                                                                                                                                                                                                                                                                                                       | 120kV chest CT, 120kV lumbar CT | AUC0.993–1.000 (95% CI:0.988–1.000)                                                                                                                                                                                        |

(Continued)

Table 2 (Continued).

| Task | Parts   | Researchers                          | Year | Purpose                                                                                                                                                                                                                                      | Algorithms Evaluated                                             | Dataset                          | Performance Metrics                                                                                                                                                                                    |
|------|---------|--------------------------------------|------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------|----------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|      | Chest   | Zhou K et al <sup>135</sup>          | 2025 | Investigate the feasibility of using DL to establish a system for vBMD prediction and OP classification based on LDCT scans                                                                                                                  | CNN                                                              | LDCT                             | OP AUC0.800(CI: Not given); Osteopenia AUC0.878 (CI: Not given)                                                                                                                                        |
|      | Chest   | Kuo DP et al <sup>136</sup>          | 2025 | Introduces an ensemble framework to leverage their complementary strengths, generating visualized and decision-transparent recommendations for DXA scans from chest LDCT.                                                                    | ViT, CNN                                                         | LDCT                             | Best performance:ViT–CNN model; Development cohort: normal AUC0.97(CI: Not given), OP AUCof 0.94(CI: Not given); Test cohort, normal AUC0.93 (CI: Not given), OP AUC0.99(CI: Not given)                |
|      | Chest   | Li Y et al <sup>137</sup>            | 2025 | Explore the feasibility and accuracy of using 100 kV low-voltage chest CT scans obtained for lung cancer screening in opportunistic osteoporosis diagnose                                                                                    | CNN                                                              | 100 kV chest CT, 120 kV chest CT | Best performance:100 kV chest CT CNN model; AUC1.000(CI: (1.000–1.000))                                                                                                                                |
|      | Chest   | Wei L et al <sup>138</sup>           | 2025 | Evaluate the diagnostic accuracy of AI in chest CT for osteoporosis/osteopenia screening in a Chinese population                                                                                                                             | Yizhun BMD AI system                                             | Chest CT                         | T10 AUC 0.84 (95% CI: 0.81–0.87), T11 AUC0.83 (95% CI: 0.80–0.86), T12 AUC0.81 (95% CI: 0.79–0.84), L1 AUC0.83 (95% CI: 0.80–0.87),                                                                    |
|      | Chest   | Zhang Y et al <sup>139</sup>         | 2026 | Develop a DL model for OP screening based on images of thoracic vertebrae obtained from 100 kV chest CT                                                                                                                                      | CNN                                                              | 100 Kv chest CT                  | AUC0.983 (CI: 0.966–1.000)                                                                                                                                                                             |
|      | Abdomen | Lim HK et al <sup>121</sup>          | 2021 | Evaluate the prediction performance of femoral OP using machine-learning analysis with radiomics features and APCT                                                                                                                           | RF                                                               | Abdomen-pelvic CT                | Training AUC0.959(95% CI 0.937–0.981); Validation AUC0.960(95% CI 0.932–0.988)                                                                                                                         |
|      | Abdomen | Tariq A et al <sup>140</sup>         | 2023 | Develop an AI model for opportunistic screening of low bone density using both contrast and non-contrast abdominopelvic CT exams                                                                                                             | CNN                                                              | Non-contrast abdominopelvic CT   | Best performance:fusion model; AUC0.86(95% CI 0.84–0.87)                                                                                                                                               |
|      | Abdomen | Yuan X et al <sup>141</sup>          | 2024 | Evaluate the performance of ML analysis based on proximal femur of abdominal CT scans in screening for abnormal bone mass in femur                                                                                                           | LR                                                               | Abdominal CT                     | Training AUC0.917 (95% CI: 0.867–0.967); Testing AUC0.963 (95% CI: 0.919–0.999)                                                                                                                        |
|      | Abdomen | Adarve-Castro A et al <sup>142</sup> | 2025 | Assess the proficiency of supervised ML techniques in discriminating between normal and abnormal BMD by leveraging clinical features and texture analysis of spinal bone tissue in patients diagnosed with primary hyperparathyroidism (PHP) | NB, LR, NN, Mixed model NN and NB                                | Abdominal CT                     | Best performance:Bayes; AUC0.916(CI: Not given)                                                                                                                                                        |
|      | Abdomen | Paukovitsch M et al <sup>143</sup>   | 2025 | Measure volumetric bone mineral density (vBMD) in clinical routine thoraco-abdominal CT using an AI-based algorithm                                                                                                                          | CNN                                                              | Thoraco-abdominal CT             | AUC0.96(95% CI: 0.93–0.98)                                                                                                                                                                             |
|      | Abdomen | Sarquis Serpa A et al <sup>144</sup> | 2026 | Develop and do multicenter validation on an algorithm that screens for OP from abdominal CTs                                                                                                                                                 | ResNet34, 2D U-Net                                               | Abdominal CT                     | Internal dataset AUC0.96 (95% CI: 0.89–1.0); External dataset AUC0.82 (95% CI: 0.74–0.89)                                                                                                              |
|      | Spine   | Yasaka K et al <sup>122</sup>        | 2020 | Investigate whether a DL model can predict the BMD of lumbar vertebrae from unenhanced abdominal CT images                                                                                                                                   | CNN                                                              | Lumbar spine CT                  | Validation dataset I: OP AUC0.965 (95% CI: 0.887–1.000), Osteopenia AUC0.955 (95% CI: 0.901–1.000); Validation dataset E: OP AUC0.970 (95% CI: 0.915–1.000), Osteopenia AUC0.903 (95% CI: 0.823–0.983) |
|      | Spine   | Tang C et al <sup>145</sup>          | 2021 | Design a novelty diagnostic method for OP screening by using the CNN                                                                                                                                                                         | U-Net, DenseNet-121                                              | Lumbar spine CT                  | AUC0.9167(CI: Not given)                                                                                                                                                                               |
|      | Spine   | Dzierżak R et al <sup>146</sup>      | 2022 | Assess the possibility of using deep DCNNs to develop an effective method for diagnosing OP based on CT images of the spine                                                                                                                  | VGG16, VGG19, MobileNetV2, Xception, ResNet50, InceptionResNetV2 | Spine CT                         | Best performance:VGG16; AUC0.985(CI: Not given)                                                                                                                                                        |
|      | Spine   | Cheng L et al <sup>147</sup>         | 2023 | Construct a combined model that integrates radiomics, clinical risk factors, and ML algorithms to diagnose OP in patients and explore its potential in clinical applications                                                                 | SVM, RF, LR                                                      | Lumbar spine CT                  | Best performance:combined model; Training AUC0.959 (95% CI: 0.9412–0.9763); Testing AUC0.910 (95% CI: 0.8690–0.9506)                                                                                   |

|  |        |                                     |      |                                                                                                                                                                                                                               |                 |                                 |                                                                                                                    |
|--|--------|-------------------------------------|------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------|---------------------------------|--------------------------------------------------------------------------------------------------------------------|
|  | Spine  | Wang J et al <sup>148</sup>         | 2024 | Develop and validate a predictive model for OP and osteopenia prediction by fusing deep transfer learning (DTL) features and classical radiomics features based on single-source dual-energy CT virtual monochromatic imaging | SVM             | Lumbar spine CT                 | Best performance:radiomics models; Training AUC0.999 (95% CI: 0.997,1.000); Testing AUC0.943 (95% CI: 0.905,0.981) |
|  | Spine  | Chen B et al <sup>149</sup>         | 2024 | Extract radiomics from CT images to screen OP in the elderly                                                                                                                                                                  | RF, SVM, kNN    | Lumbar spine CT                 | Best performance:RF; Training AUC0.925 (95% CI: 0.922–0.939); Test AUC0.826 (95% CI: 0.809–0.836)                  |
|  | Spine  | Liu J et al <sup>150</sup>          | 2024 | Proposed a transformer-enhanced DL framework to automatically acquire features from the entire lumbar vertebral body and the cancellous compartment to develop radiomics models for OP screening in routine CT                | RF, Transformer | MDCT                            | Training AUC0.95(95% CI: 0.91–0.99), Test AUC0.97 (95% CI: 0.93–1.00)                                              |
|  | Spine  | Küçükçiloğlu Y et al <sup>151</sup> | 2024 | Develop unimodal and multimodal deep-learning-based diagnostic models to predict bone mineral loss of the lumbar vertebrae using MRI and CT imaging                                                                           | CNN             | Lumbar spine MRI or CT          | Best performance:multimodal model; Accuracy0.9890 (95% CI: 0.984–0.994)                                            |
|  | Others | Xu Y et al <sup>152</sup>           | 2013 | Propose a novel method based on machine-learning method performed on micro-CT images to diagnose OP                                                                                                                           | SVM, kNN        | Micro-CT                        | Precision1.00(CI: Not given)                                                                                       |
|  | Others | Namatevs I et al <sup>123</sup>     | 2023 | Examine the capabilities of deep DCNNs for diagnosing OP through cone-beam computed tomography (CBCT) scans of the mandible                                                                                                   | ResNet-101      | CBCT                            | Training Accuracy0.9885(CI: Not given); Validation Accuracy0.9399(CI: Not given)                                   |
|  | Others | Sebro R et al <sup>153</sup>        | 2024 | Evaluate whether the CT attenuation of bones seen on shoulder CT scans could be used to predict low BMD                                                                                                                       | SVM, kNN        | Shoulder CT                     | Best performance:SVM; AUC0.857(CI: Not given)                                                                      |
|  | Others | Du C et al <sup>154</sup>           | 2025 | Establish an automated OP detection model based on low-dose abdominal CT                                                                                                                                                      | RF              | Proximal femur images from LDCT | OP AUC0.960 (95% CI: 0.913–1.000); Osteopenia, AUC0.828 (95% CI: 0.747–0.909).                                     |

(Continued)

Table 2 (Continued).

| Task                                  | Parts   | Researchers                      | Year | Purpose                                                                                                                                                                                                                                    | Algorithms Evaluated                                      | Dataset                         | Performance Metrics                                                                                                                                      |
|---------------------------------------|---------|----------------------------------|------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------|---------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------|
| BMD Envaluate                         | Chest   | Pan Y et al <sup>155</sup>       | 2020 | Develop a DL -based system to automatically measure BMD for opportunistic OP screening using low-dose chest computed tomography (LDCT) scans                                                                                               | U-net                                                     | LDCT                            | OP AUC0.927(95% CI: 0.896–0.951); Low BMD AUC0.942(95% CI: 0.914–0.964)                                                                                  |
|                                       | Chest   | Pan Y et al <sup>156</sup>       | 2023 | Develop and evaluate a fully automated method based on DL and phantomless internal calibration for BMD measurement and opportunistic low BMD (osteopenia and OP) screening using chest LDCT scans                                          | CNN                                                       | Chest LDCT                      | VCI: OP AUC0.964(95% CI:0.947–0.976); Low BMD AUC0.943(95% CI:0.923–0.959); VCII: OP AUC0.968 (95% CI:0.940–0.985); Low BMD AUC0.984(95% CI:0.962–0.995) |
|                                       | Chest   | Hathaway QA et al <sup>157</sup> | 2025 | Determine whether nnU-net can measure 3D vertebral body BMD across consistently imaged thoracic levels (T1–T10) at any conventional, noncontract chest CT examination.                                                                     | nnU-net                                                   | Chest CT                        | Best performance:3D vertebral BMD with FRAXnb; AUC =0.82(CI: Not given)                                                                                  |
|                                       | Abdomen | Oh J et al <sup>141</sup>        | 2024 | Train DL models to localize the center axial slice from the CT scan of the torso, segment the vertebral bone, and estimate BMD for the top four lumbar vertebrae                                                                           | U-Net, DenseNet169                                        | Abdominal CT                    | MAE0.066, Pearson's r0.907 (P<0.001), R2 of 0.78(CI: Not given)                                                                                          |
|                                       | Spine   | Fang Y et al <sup>158</sup>      | 2021 | Develop a fully automatic method based on deep DCNN for vertebral body segmentation and BMD calculation in CT images                                                                                                                       | U-Net, DenseNet-121                                       | Lumbar spine CT                 | Minimum average dice coefficients 0.823 (95% CI: 0.815–0.831), 0.786 (95% CI: 0.773–0.799), and 0.782 (95% CI: 0.769–0.797)                              |
|                                       | QCT     | Oh S et al <sup>159</sup>        | 2024 | Describe a new DL algorithm for vertebral segmentation and localizing of L1 and L2 vertebrae in routine clinical chest, abdomen, and lumbar spine CT scans and automatically draw a ROI from each vertebra to calculate volumetric QCT BMD | DL, U-Net                                                 | QCT                             | Low-BMD AUC0.847(CI: Not given); OP AUC0.770 (CI: Not given)                                                                                             |
|                                       | QCT     | Li Y et al <sup>160</sup>        | 2025 | Evaluate the diagnostic efficacy of AI-based automated BMD measurement system for opportunistic OP screening using lumbar DECT scans                                                                                                       | 3D RetinaNet, U-Net                                       | DECT, QCT                       | OP AUC0.979(95% CI: 0.935–0.997); Low BMD AUC0.980 (95% CI: 0.935–0.997)                                                                                 |
|                                       | Others  | Schmidt D et al <sup>161</sup>   | 2022 | Evaluate a fully automated DL -based method for lumbar vertebral segmentation and measurement of vertebral volumetric trabecular attenuation values                                                                                        | CNN                                                       | Non-contrast-enhanced CT        | Mean L1 attenuation value 140 HU ± 54(CI: Not given)                                                                                                     |
| Fracture Detection or Risk Prediction | Chest   | Tomita N et al <sup>120</sup>    | 2018 | Developed a system to detect osteoporotic vertebral fractures (OVFs) on CT exams                                                                                                                                                           | LSTM, ResNet34                                            | Chest CT, abdomen CT, pelvis CT | Accuracy0.892(CI: Not given)                                                                                                                             |
|                                       | Spine   | Biamonte E et al <sup>162</sup>  | 2022 | Evaluate radiomics features on CT images of the lumbar spine in subjects with or without fragility vertebral fractures                                                                                                                     | LSVM                                                      | Lumbar spine CT                 | Training AUC0.839(CI: Not given); Testing AUC0.789 (CI: Not given)                                                                                       |
|                                       | Spine   | Zhang J et al <sup>163</sup>     | 2024 | Develop and validate a predictive model for OVFs risk by integrating demographic, BMD, CT imaging, and DL radiomics features from CT images                                                                                                | CNN, SVM, KNN, LR, RF, Extratrees, XGBoost, LightGBM, MLP | Spine CT                        | Best performance:fusion model; Training AUC0.980 (95% CI: 0.959–1.000); Test AUC0.971 (95% CI: 0.928–1.000)                                              |

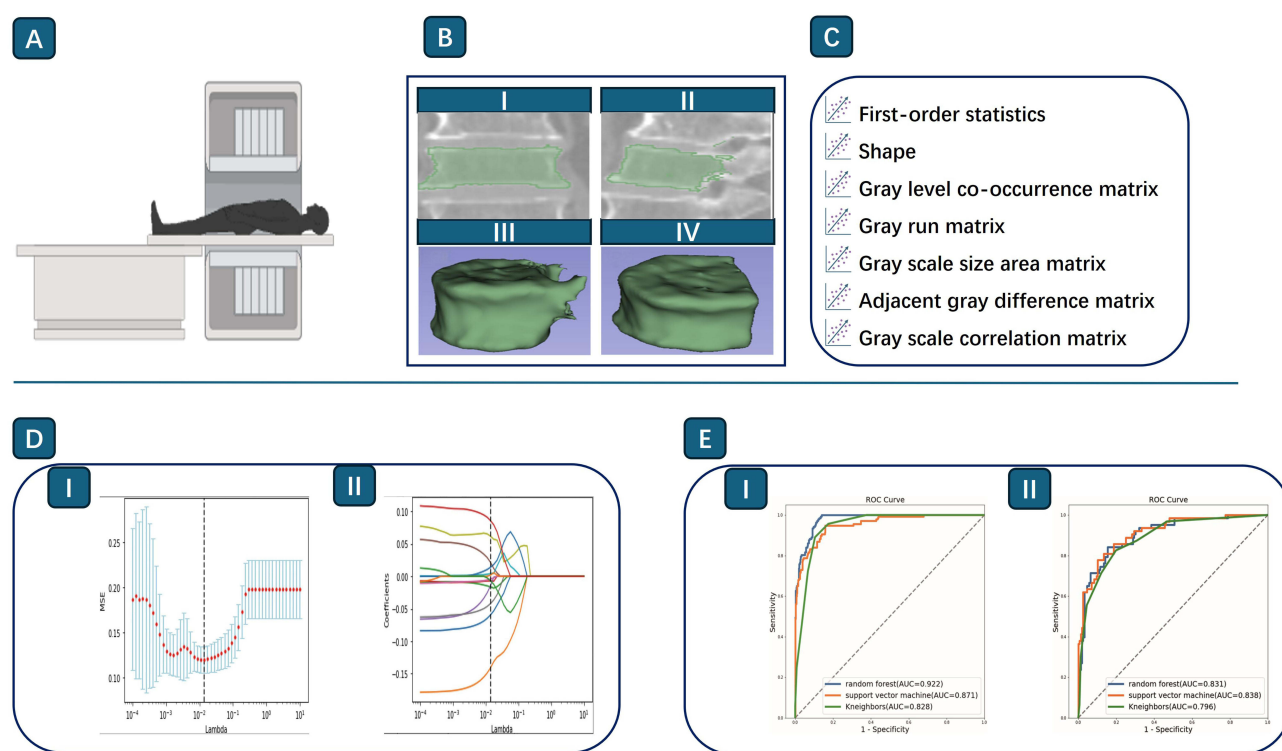
|                                              |         |                                |      |                                                                                                                                                                                                                 |                                                                                                  |                  |                                                                                                   |
|----------------------------------------------|---------|--------------------------------|------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|------------------|---------------------------------------------------------------------------------------------------|
| Bone Quality or Microarchitecture Prediction | Spine   | Yoshida K et al <sup>164</sup> | 2023 | Evaluate the feasibility of using DL with a CNN for predicting BMD and bone microarchitecture from conventional CT images                                                                                       | ResNet50                                                                                         | Lumbar spine CT  | Osteopenia AUC0.921(95% CI: 0.793–0.973); OP AUC0.969 (95% CI: 0.872–0.993)                       |
|                                              | QCT     | Zhang K et al <sup>124</sup>   | 2024 | Develop a novel multi-domain diagnostic method for learning domain-invariant features from CT images, enhancing OP analysis generalization across varied CT regions and accommodating dose or device variations | DeepmdQCT, AlexNet, ResNet-50, ResNet-101, DenseNet-101, ShuffleNet, EfficientNet, DSNet, DLGNet | QCT              | Dose domain image Accuracy0.910(CI: Not given); Device domain images Accuracy0.905(CI: Not given) |
| Technical Development                        | Spine   | Feng E et al <sup>165</sup>    | 2025 | Develop an AI model for automatically detecting Hounsfield unit (HU) values at the L1 vertebra in preoperative thoracolumbar CT scans                                                                           | nnU-Net                                                                                          | Thoracolumbar CT | High Dice coefficient 0.91(CI: Not given)                                                         |
|                                              | Abdomen | Song J et al <sup>166</sup>    | 2026 | Assess whether integrating volumetric vertebral and body composition features obtained from DL segmentation of CT images enhances the prediction of BMD and the classification of OP                            | 3DnnU-Net                                                                                        | Abdomen CT       | AUC0.95(CI: Not given)                                                                            |
|                                              | Spine   | Hiyama A et al <sup>167</sup>  | 2026 | Evaluate the utility of L1–L4 average HU values from lumbar spine CT in predicting OP using multiple ML models                                                                                                  | LR, RF, SVM, KNN, NB, DT, XGBoost                                                                | Lumbar spine CT  | Best performance: KNN; AUC0.705(CI: 0.659–0.751)                                                  |

**Abbreviations:** AI, artificial intelligence; BMD, bone mineral density; vBMD, volumetric bone mineral density; OP, osteoporosis; OVf, osteoporotic vertebral fracture; CT, computed tomography; QCT, quantitative computed tomography; DXA, dual-energy X-ray absorptiometry; ROI, region of interest; HU, Hounsfield unit; CNN, convolutional neural network; DCNN, deep convolutional neural network; U-Net, U-shaped convolutional neural network; DenseNet, densely connected convolutional network; ResNet, residual neural network; VB-Net, volumetric bone network; LSTM, long short-term memory network; RF, random forest; RFC, random forest classifier; SVM, support vector machine; KNN, k-nearest neighbors; KNNC, k-nearest neighbors classifier; LR, logistic regression; MLP, multilayer perceptron; VMC, virtual monochromatic imaging; AUC, area under the receiver operating characteristic curve; ACC, accuracy; SEN, sensitivity; SPE, specificity; MAE, mean absolute error; RMSE, root mean square error; CI, confidence interval.

- (2) ROI Segmentation: Radiologists manually delineate volumetric regions of interest (VOIs) in vertebral bodies using 3D Slicer, excluding cortical bone and venous plexuses.
- (3) Feature Extraction: First-order, shape, and texture features are extracted via PyRadiomics, while transfer learning features are derived from a pretrained ResNet50 model.
- (4) Feature Selection: Redundant features are removed through reproducibility testing, Spearman correlation, and LASSO regression.
- (5) Application: Selected features are used to train a two-level SVM classifier, which distinguishes OP from non-OP cases and further differentiates osteopenia from normal BMD (Figure 2).<sup>148–150</sup>

Studies such as those by Liu et al and Du et al have successfully applied radiomics to extract lumbar vertebral features from abdominal CT scans,<sup>154,170</sup> enabling quantitative and high-dimensional OP diagnosis. Radiomics allows for the integration of imaging, molecular biology, pathology, and data science, and is likely to represent a promising analytical direction for future CT-based OP research.

Oromaxillofacial CT is another clinically common modality, offering practical potential for OP diagnosis. In 2023, Park et al developed a DL-based QCBCT-Net model to analyze quantitative cone-beam CT (QCBCT), achieving a root mean square error (RMSE) of 83.41 mg/cm<sup>3</sup>, indicating high precision.<sup>171</sup> Similarly, Namatevs et al created a DCNN-based AI tool for detecting OP from CBCT images, reporting a classification accuracy of 93.99%.<sup>123</sup> These findings highlight the feasibility of using oromaxillofacial CT for OP assessment and suggest a promising research direction for automated dental OP screening. Integration of AI with qCT to improve its diagnostic power is another emerging trend. In 2024, Oh et al developed



**Figure 2** A schematic workflow of CT-based OP diagnosis using radiomics, illustrated using lumbar spine CT as an example. (Adapted and redrawn based on Figures 2,3 and 4 from Cui Z et al, "Application of radiomics model based on lumbar computed tomography in diagnosis of elderly osteoporosis," with permission).<sup>148–150</sup> **(A)** Image Acquisition: Use CT to acquire the image of the target part of the body. **(B)** ROI Segmentation: ROI are delineated by radiologists using 3D Slicer. BI. Segmentation of image coronal section. BII. Segmentation of sagittal slice of image. BIII. 3D model construction. IV. Model construction processing. **(C)** Feature Extraction: Features are extracted via PyRadiomics. **(D)** Feature Selection: Useful features are selected by LASSO regression. DI. Best lambda value. A ten-fold cross-validation procedure was applied to identify the optimal lambda parameter. The y-axis shows the mean squared error (MSE), while the x-axis represents the lambda sequence. Vertical dashed lines denote the optimal lambda value, defined as the lambda yielding the minimum average MSE regardless of its spread. DII. LASSO coefficient profile of radiomic features. Coefficient trajectories of fourteen radiomic features are plotted against the lambda sequence. The vertical dashed lines indicate the optimal lambda value. **(E)** Application: Selected features are used to train models. EI. ROC curves of training set. EII. ROC curves of test set.

a DL model based on Python (v3.8.5) and TensorFlow-GPU (v2.4.0) to automatically measure BMD from qCT images. Using DXA as a reference, the model achieved AUCs of 0.847 and 0.770 for distinguishing low BMD and OP, respectively.<sup>159</sup> In the same year, Zhang et al addressed the issue of inter-device variability by developing a DeepmdQCT model built on a ResNet architecture with a comprehensive attention-guided module (CAGM) that integrates both global and local features. The model achieved average accuracies of 91% (normal dose) and 90.5% (low dose), maintaining consistent performance across Philips and Siemens imaging systems—demonstrating strong generalizability.<sup>124</sup>

From the above studies, it can be observed that CT can provide three-dimensional structural information of the measurement sites,<sup>172</sup> and when combined with CNN-based models, it enables accurate bone mineral density analysis.<sup>126</sup> However, compared with X-ray imaging, CT is associated with higher costs and greater radiation exposure.<sup>173</sup> Therefore, although CT-based approaches offer advantages in terms of diagnostic accuracy, their widespread use in large-scale or initial screening is limited.

## MRI and Ultrasound Imaging

Compared with X-ray and CT imaging, studies that combine MRI or ultrasound (US) with AI for OP diagnosis remain relatively limited.

In the MRI-based research field, one of the earliest studies was conducted by Ferizi et al, who applied multiple ML algorithms to predict the risk of OF using MRI data. They evaluated 15 ML models, including SVM and k-NN, while incorporating patient-level features such as age, body weight, and height. Among the tested models, RUS-boosted trees, logistic regression(LR), and linear discriminant analysis (LDA) achieved the best predictive performance.<sup>174</sup> More recently, in 2024, Kūçukçiloğlu et al integrated MRI and CT data for multimodal AI training. They developed four distinct models: (1) a single-modality MRI model, (2) a single-modality CT model, (3) a combined single-modality model trained on merged MRI–CT data, and (4) a multimodal model treating MRI and CT as separate inputs. The multimodal model achieved the highest performance, with a balanced accuracy of 98.90% on MRI–CT datasets and similarly strong accuracy in patient-level validation.<sup>151</sup>

In the US-based domain, Vogl et al (2019) introduced an AI-assisted framework using low-frequency guided waves to assess acoustic parameters of the tibia, followed by SVM classification for OP diagnosis.<sup>175</sup> However, because low-frequency guided waves are not standard in clinical ultrasonography, the practical applicability of this method remains uncertain. In the same year, Mohanty et al estimated cortical bone microstructural parameters from US frequency-dependent attenuation, combined with 2D finite-difference time-domain (FDTD) simulations to generate simplified cortical models. These were refined using CT-derived structural data and subsequently used to train an ANN to predict pore diameter, pore density, and porosity. The ANN demonstrated high prediction accuracy, suggesting that AI-augmented US could become a promising tool for OP assessment.<sup>176</sup> In recent years, Ferguson HE proposed combining CNNs with US backscatter to assess BMD, achieving promising results. This further demonstrates the feasibility of US applications in the field of OP.<sup>177</sup>

Overall, the relatively limited use of MRI and US in AI-assisted OP diagnosis may be attributed to their lower prevalence in orthopedic imaging and the inherent limitations of these modalities for bone evaluation,<sup>178, 179</sup> Nonetheless, the ability of AI to extract subtle diagnostic information from MRI and US data suggests that AI-enhanced multimodal imaging could represent a valuable future direction in OP research.

Although imaging-based AI has demonstrated strong performance in identifying OP and related fractures, these approaches primarily focus on phenotypic manifestations of the disease. To understand the mechanisms of the disease, researchers have increasingly turned to bioinformatics and genetics. By integrating AI with molecular-level data, it has become possible to explore the genetic basis and biological pathways underlying OP.

## Identification of Diagnostic Genes

In addition to imaging-based diagnosis, AI and ML models have also been applied to gene-based diagnostic prediction in OP. In these studies, selected genes or gene-expression profiles are commonly used as input features for classification models aimed at distinguishing osteoporotic from non-osteoporotic populations.<sup>180</sup> Unlike studies focusing on molecular mechanisms, these approaches primarily aim to develop predictive or classification models capable of distinguishing osteoporotic

individuals from non-osteoporotic populations. For example, Hu et al identified five key DEGs—CCR1, CD33, HCK, LILRB2, and CYBB—through bioinformatics analysis and subsequently constructed an SVM classification model based on these genes, achieving accurate classification and prediction of OP samples.<sup>50</sup> Similarly, Ding et al and Zheng et al adopted comparable approaches: Ding integrated gene expression profiles with clinical features using both MLR and ANN models,<sup>181</sup> while Zheng systematically evaluated diagnostic performance across different gene combinations to determine the optimal gene subset.<sup>43</sup> Building upon this concept, Lin et al focused on the effect of smoking on BMD. Using bioinformatics methods, they identified gene modules associated with BMD in both smokers and non-smokers and selected ten shared genes (including TNS4 and IRF2) to construct an SVM–RFE model. This model effectively distinguished high- and low-BMD individuals in both groups, achieving an AUC > 0.9, thereby demonstrating strong potential for identifying individuals at high risk of OP.<sup>182</sup> These studies demonstrate the potential of combining bioinformatics and AI for OP diagnosis and risk stratification. Broader applications of AI in molecular mechanism exploration and biomarker discovery are discussed further in [Artificial Intelligence in Genetic and Molecular Studies of Osteoporosis](#).

## Non-Imaging AI-Based Diagnostic Technologies

Beyond imaging and genomics, several novel AI-based diagnostic methods have been proposed. For instance, Yang et al suggested that surface-enhanced Raman scattering (SERS) spectral features from blood samples could be analyzed using SVM to construct an OP diagnostic model.<sup>183</sup> Although this represents a promising alternative approach, the current mainstream of AI-assisted OP diagnosis still relies predominantly on medical imaging. In 2026, Liang Q et al conducted a study that did not utilize any medical imaging, relying solely on ordinary facial photographs to diagnose OP, thereby further facilitating its detection.<sup>184</sup> Nevertheless, these emerging diagnostic modalities offer broad opportunities for future exploration and may complement imaging-based AI strategies in clinical practice.

## Artificial Intelligence in Genetic and Molecular Studies of Osteoporosis

Bioinformatics is an interdisciplinary field that applies computational, mathematical, and statistical methods to analyze biological data, such as genomic and proteomic information.<sup>185</sup> It has been extensively used to explore disease-associated genes and proteins, providing potential diagnostic and therapeutic targets for various diseases. With the rapid development of AI, bioinformatics has gradually merged with AI technologies, a trend that has revolutionized biomedical research. The integration of AI and bioinformatics has already yielded promising results in gene sequence alignment and annotation, non-coding RNA prediction, protein folding analysis, protein–protein interaction modeling, and drug discovery and development.<sup>186,187,188</sup> In the field of OP, beyond diagnostic applications,<sup>23</sup> AI has increasingly been utilized to investigate the molecular mechanisms underlying OP, for example, therapeutic targets,<sup>189</sup> as well as for the identification of genes shared with other diseases.<sup>190</sup> Recently, AI techniques have been increasingly incorporated into these studies, significantly improving efficiency and analytical depth.

## AI-Based Biomarker Identification for Osteoporosis

Bioinformatics is frequently applied to identify diagnostic and therapeutic targets for various diseases. In OP research, several studies have utilized bioinformatics to uncover such targets. For example, Zhang L et al identified the methylation biomarker MAP3K5 as associated with OP and validated its potential therapeutic value through immune cell infiltration analysis.<sup>191</sup> With the advancement of AI, integrating intelligent algorithms into bioinformatics pipelines has become an emerging trend, greatly accelerating research progress. As early as 2010, Guan Y et al proposed combining functional genomics with AI for disease-gene prediction. They constructed functional relationship networks and employed SVM to predict phenotype associations, successfully identifying BMD-related genes such as Timp2 and Abcg8. Experimental validation in mice confirmed that knocking out these genes significantly reduced BMD, demonstrating the feasibility of AI-assisted genomic analysis in OP research.<sup>192</sup> Feng ZW et al (2024) followed a similar workflow.<sup>193</sup> In 2019, Yang C et al identified DHTKD1, OSTF1, and GPR116 as key genes associated with OP, and validated their relevance using multiple ML models, including SVM, decision tree, and RF, though without animal validation.<sup>194</sup> In addition, Long SW et al used a RF model to identify five ferroptosis-related hub genes (CP, FLT3, HAMP, HMOX1, SLC2A3), constructed diagnostic models, and confirmed their roles experimentally, establishing these genes as potential biomarkers for OP.<sup>195</sup>

Some other studies combined traditional bioinformatics-based gene screening with AI validation, which were discussed earlier and will not be elaborated here. Overall, AI can play a major supporting role in identifying OP-related diagnostic and therapeutic biomarkers, and future studies are expected to expand in this promising direction.

## AI-Driven Molecular Mechanism Exploration

Beyond identifying diagnostic biomarkers, AI has been increasingly applied to elucidate the molecular mechanisms underlying OP by analyzing gene networks and signaling pathways. Understanding the molecular mechanisms underlying disease pathogenesis is crucial for developing targeted treatments. In OP, bioinformatics-based molecular analyses have long been conducted, but the introduction of AI has injected new vitality into this research area. Xiao KW et al investigated how monocytes affect BMD through the ribonucleoprotein complex biosynthesis pathway. After identifying key genes via gene set enrichment analysis (GSEA), they built an elastic net regression model to predict BMD. It is an early but imperfect attempt to apply AI to mechanistic studies of OP.<sup>196</sup> Subsequent research has increasingly focused on immune-related mechanisms. Hao S et al used LASSO and mSVM-RFE algorithms to identify CCR5 and IAPP as key BMD-related immune genes, validating their roles experimentally—the first study to combine bioinformatics and ML to identify immune-related OP genes.<sup>197</sup>

Similarly, Chen L et al (2024) analyzed postmenopausal OP (PMOP) using LASSO and RF models, identifying PYGM and POMP as regulators of immune response and proteolysis, marking their first report in PMOP.<sup>198</sup> Zhang B et al further revealed that neutrophils modulate BMD via inflammatory cytokine secretion.<sup>199</sup> Li J et al (2025) focused on FBXW4, applying ML to identify its co-expressed hub genes and discovering that FBXW4 may regulate OP progression via antiviral defense, cytokine production, and immune response modulation. Beyond immunity, other mechanisms have been explored.<sup>200</sup> Wang X et al used ML to identify ferroptosis-related molecular subtypes in diabetic osteoporosis (DO) and identified IDH1 as a key gene.<sup>201</sup> Feng Z et al<sup>202</sup> Bi K et al<sup>203</sup> and Li S et al<sup>204</sup> focused on pyrimidine metabolism genes (PyMGs), cellular senescence, mitochondrial biomarkers, and smoking-related genes, broadening the mechanistic landscape of OP. In 2025, Zeng HB et al also focused on the role of mitochondrial dysfunction in OP, identifying three key genes—ALAS1, HSPB1, and VPS35. They further proposed that overexpression of VPS35 inhibits osteoblast differentiation by suppressing the ERK/PI3K/AKT signaling pathway, a process regulated by miR-142-5p.<sup>205</sup> A particularly novel study by Zhang X et al combined network toxicology, ML, and molecular docking to examine plasticizer-induced OP. Using 113 ML models (LASSO, SVM, RF, etc.), they identified core genes linking plasticizer exposure to OP, pioneering research into the toxicological mechanisms of OP. This integrative framework—combining target screening, model optimization, and molecular validation—offers new perspectives for future studies.<sup>51</sup>

## AI-Based Cross-Disease Genetic Analysis

A growing number of studies have revealed shared genes between OP and other diseases, such as atherosclerosis (Mishra BH et al)<sup>206</sup> and type 2 diabetes mellitus (T2DM) (Du A et al).<sup>207</sup> Investigating shared genetic profiles can uncover common disease mechanisms, reveal potential therapeutic targets, and facilitate personalized and multi-disease intervention strategies. The incorporation of AI has accelerated progress in this area. For example, Zhao R et al focused on OP and T2DM. Using LR, cross-analysis, and RF algorithms, they identified three hub genes.<sup>208</sup> In another study, Liu J et al applied XGBoost to identify pyroptosis- and crosstalk-related hub genes between periodontitis and OP providing insights into shared mechanisms.<sup>209</sup> Yang J et al examined chronic HBV infection and OP using differential gene expression and LASSO regression, identifying three shared genes—USP10, ERL1, and ECM1—which may assist in the diagnosis and management of HBV-related OP.<sup>52</sup> Similarly, Xu G et al explored inflammatory bowel disease (IBD) and OP, using LASSO combined with NETs-related gene analysis to identify HDAC6, IL-8, and PPIF as diagnostic genes. Their study revealed potential neutrophil extracellular trap (NET) involvement in the connection between IBD and OP.<sup>210</sup> In 2026, Tang H et al employed ML to investigate shared genes between OP and chronic kidney disease (CKD), ultimately identifying four genes—FAM184A, NFKBIA, RP2, and HIRA—providing a molecular-level explanation for the high prevalence of OP in CKD patients.<sup>211</sup> Collectively, these findings highlight AI's ability to uncover shared molecular mechanisms among diseases. However, most current research focuses on pairwise disease associations; future work should aim to investigate multi-disease gene networks, ultimately enabling multi-target therapeutic strategies.

Beyond these primary areas, the cross-disease genetic analysis has also been explored in drug discovery. For instance, Yang X et al identified shared genes between OP and sarcopenia using ML models and subsequently employed the CMap database to predict potential therapeutic compounds.<sup>212</sup> Zhang C et al investigated the shared genes between peri-implantitis and OP, identifying three common DEGs, namely ALDH1A3, MGP, and CYBB. They further explored potential drug targets associated with these hub genes.<sup>213</sup> In the same year, Lv Y et al applied ML models, including LASSO, to identify five shared genes between sarcopenia and OP—APOC1, ENPP5, FBXL22, IRS1, and PAQR4—revealing the common molecular mechanisms underlying these two age-related degenerative conditions.<sup>214</sup> These applications will be discussed in detail in the following section on AI-assisted treatment of OP.

Insights gained from AI-driven genetic and molecular analyses not only enhance the understanding of OP pathogenesis but also provide a foundation for individualized risk assessment. Building upon these mechanistic findings, AI has been further applied to predict disease risk, stratify patient populations, and estimate future clinical outcomes.

## AI in Osteoporosis Prediction and Risk Stratification

Accurate identification of risk factors and prognostic prediction for OP are crucial for achieving precise diagnosis and stratified management.<sup>212</sup> These approaches not only optimize the use of medical resources but also spare patients from unnecessary examinations and treatments. In recent years, AI has been increasingly adopted to identify disease risk factors and predict clinical outcomes.<sup>215, 216</sup> For instance, Doppalapudi S et al employed ANNs, recurrent neural networks (RNNs), and CNNs to analyze demographic data, tumor site and morphology, disease stage, and treatment regimens, successfully predicting lung cancer prognosis.<sup>217</sup> Similarly, multiple studies have integrated AI into OP risk assessment and outcome prediction, demonstrating its growing potential in clinical decision support.

## ML Applications

Since research on OP-related risk factors often involves structured data (eg., age, body mass index(BMI), blood markers), ML—rather than DL—is particularly well suited for this task.<sup>218</sup> PMOP has long been a major public health issue. Identifying predictive factors can facilitate early prevention. As early as 2013, Yoo TK et al applied several ML algorithms—SVM, RF, ANN, and LR—to analyze patient characteristics. They identified age, height, weight, BMI, duration of menopause, breastfeeding duration, estrogen therapy, hyperlipidemia, hypertension, osteoarthritis, and diabetes as key risk factors. Among these models, SVM achieved the best performance on both training and validation datasets.<sup>219</sup> Similarly, Jin W et al used EHRs from female patients, combining four feature-selection methods and eight ML algorithms to predict hip OP. Their final model achieved an external validation sensitivity of 0.775.<sup>220</sup> Beyond predicting OP, ML has been applied to OF prediction. In 2021, de Vries BCS et al compared multiple models—including random survival forest (RSF)—for predicting major osteoporotic fractures (MOFs), finding Cox regression performed best.<sup>221</sup>

For individuals below the recommended age for DXA screening, Park HW et al (2021) used XGBoost, LR, and multilayer perceptron (MLP) models to identify OP risk factors. The XGBoost model outperformed traditional risk assessment tools, with age, weight-related variables (BMI, weight, obesity), serum alkaline phosphatase (ALP), systolic blood pressure (SBP), blood urea nitrogen (BUN), and alcohol consumption identified as major predictors.<sup>222</sup> ML has also proven valuable for patients with chronic diseases, who are at higher risk of secondary OP. Peng Y et al used clinical data—including genetic markers—from elderly patients with cardiovascular risk to predict OP. Among four ML models compared with LR, the latter performed best, underscoring the value of model benchmarking in medical AI research.<sup>223</sup> Similar methods were employed by Hsu CT,<sup>224</sup> Wei Q,<sup>225</sup> and Yu X,<sup>226</sup> focusing on patients with CKD and type 2 diabetes, respectively. ML has additionally been applied to postoperative outcome prediction in OP treatment. Klemm C et al used neural networks and RF to predict surgical revision risks, identifying female sex, BMI > 35 kg/m<sup>2</sup>, age > 70 years, ASA score ≥ 3, and T-score as the strongest predictors.<sup>227</sup> Beyond traditional demographic and clinical variables, some studies have incorporated imaging-derived features into ML models. For instance, Liu L et al developed a three-tier ML model for OP diagnosis: Tier 1 used only demographic features, Tier 2 incorporated clinical data, and Tier 3 added CT imaging features. The third-tier model performed best, demonstrating the added value of imaging information.<sup>228</sup> Recent studies have shifted toward bone microarchitecture and geometric analysis. Mateo J et al (2025) used trabecular bone score (TBS) and 3D-DXA-derived microstructural and geometric parameters to predict fracture risk in elderly

women, achieving an internal validation accuracy of  $89.24\% \pm 0.52\%$  with an RF model.<sup>229</sup> Similar work by Quagliato L et al supported these findings.<sup>115</sup> Incorporating imaging data has further expanded predictive capabilities. Sebro R analyzed CT attenuation values from multiple skeletal regions on shoulder and chest CT to predict OP,<sup>230,231</sup> Huang CB evaluated psoas muscle index (PMI) as a predictor,<sup>232</sup> and Zhang J et al used DL-extracted radiomic features for OP prediction.<sup>163</sup> Other studies have explored unconventional biomarkers. For instance, Kang SJ proposed metal element profiling in hair samples as a potential OP predictor.<sup>233</sup> A similar study was conducted by Huang W et al. However, their study focused on heavy metals in the urinary tract.<sup>234</sup> Such innovative approaches may reveal new biomarkers for early OP detection in the future.

## Other Artificial Intelligence Techniques

Beyond classical ML, other AI approaches have also been employed in OP risk prediction, particularly ANNs. As early as 1999, Queralto JM et al used ANN models to predict bone loss in postmenopausal women, based on plasma estrogen, osteocalcin, parathyroid hormone (PTH), and urinary calcium and hydroxyproline.<sup>235</sup> Similarly, Sadatsafavi M et al (2005) used ANN models with age, body weight, menopausal age, corticosteroid and estrogen use, parity, menarche age, height, exercise, and smoking as inputs to predict BMD, achieving significantly higher AUROC than regression models.<sup>236</sup> Several comparative studies have evaluated ANN versus ML models. Yoo TK et al<sup>219</sup> Xu R et al<sup>54</sup> and Shim JG et al<sup>44</sup> all conducted such comparisons; while the first two found ML models superior, the latter reported ANN outperformed others—differences likely due to variations in input features and dataset characteristics. Interestingly, in contrast to the studies that primarily focused on elderly populations, the study by Jiang et al applied an ANN to investigate metabolic bone disease in neonates. Their work explored the effects of various prenatal and postnatal factors on disease risk and demonstrated that the ANN was effective in identifying key risk factors for neonatal OP. Among these, extremely low birth weight and antenatal magnesium sulfate exposure emerged as the two most significant contributors.<sup>237</sup> More recently, DL methods have gained traction. Suh B et al (2023) developed an interpretable DL model based on clinical features to screen for OP. Key predictors included sex, age, BMI, arm circumference, obesity prevalence, and socioeconomic status, with DL showing markedly higher accuracy than conventional models.<sup>238</sup> Similar DL-based approaches were reported by Hung WC<sup>239</sup> and Cho SW,<sup>24</sup> reinforcing DL's growing importance in OP risk assessment. In addition, Cho et al employed DL to extract imaging features from spinal radiographs and performed an analysis of spinal age. Their findings revealed a significant association between spinal age and OP. Although the study primarily focused on elderly patients with fractures, this observation nevertheless highlights the potential of DL-based approaches to capture osteoporosis-related imaging features from radiographs.<sup>240</sup> Compared with the previous study, Tang J et al not only extracted imaging features using a DL model but also integrated patients' clinical characteristics to construct a multimodal model. This multimodal model achieved an AUC of 0.975.<sup>241</sup> As DL techniques evolve, their applications in OP prediction and prevention are expected to become increasingly refined and clinically impactful.

Risk prediction and stratification represent a critical step toward precision medicine in OP. Beyond identifying individuals at high risk, AI has also been explored as a tool to support therapeutic decision-making, drug discovery, and treatment outcome prediction, thereby extending its role from risk assessment to clinical intervention.

## AI in the Treatment of Osteoporosis

Compared with its extensive applications in OP diagnosis, risk factor identification, prognosis prediction, and bioinformatics research, studies on the use of AI in OP treatment remain relatively limited. Current AI-assisted therapeutic research mainly focuses on identifying key genes within osteoporotic molecular pathways to discover potential therapeutic compounds,<sup>55</sup> optimizing and personalizing existing treatment strategies, and evaluating postoperative or long-term treatment outcomes.<sup>242</sup>

In drug discovery, bioinformatics-based AI approaches are most widely applied. For instance, Yang X et al utilized LASSO, SVM-RFE, and RF algorithms to identify CHST3, PGBD5, and SLIT2 as comorbidity-related diagnostic genes shared between OP and sarcopenia. These genes were subsequently analyzed using the CMap database to predict potential therapeutic agents targeting both diseases.<sup>55</sup> Similarly, Long SW et al employed an RF model to identify ferroptosis-related biomarkers from the GEO database and applied molecular docking to screen for small-molecule

compounds with therapeutic potential against OP.<sup>195</sup> A recent study in this field by Li Q et al utilized Chemprop to develop a predictive model for cathepsin K (CTSK) inhibition. Experimental validation identified three compounds, including quercetin, that exhibited the strongest concentration-dependent CTSK inhibitory effects. These compounds represent promising candidate drugs for the treatment of osteoporosis.<sup>243</sup> Beyond novel target discovery, AI has also been used to explore repurposing of existing drugs for OP treatment. For example, Hung TNK et al investigated the pharmacological interactions between medications used for Paget's disease and OP, offering new insights into cross-disease drug repositioning.<sup>42</sup> In addition to identifying new or repurposed drugs, another important direction is using AI to recommend optimal treatment regimens from existing therapies. Bonaccorsi G et al developed a ML-based clinical decision system that integrates patient data to recommend personalized treatment strategies. The model demonstrated high predictive accuracy across various treatment-related decisions—including bone-protective therapy, vitamin D supplementation, and calcium administration—achieving accuracies approaching 90% in some cases.<sup>244</sup> Lin YT et al (2022) collected a comprehensive dataset of 33 clinical variables, encompassing patient demographics (age, sex, height, weight), laboratory data (serum calcium, phosphate levels), and medications (eg., alendronate, raloxifene, teriparatide), to train four ML models. These models successfully predicted therapeutic outcomes and assisted clinicians in adjusting individualized treatment plans.<sup>242</sup> Similarly, in 2026, Sugawara Y et al utilized patients' clinical and imaging features to train five ML models to assist clinical drug decision-making. They found that the LightGBM model among the five achieved an accuracy above 0.9, highlighting the potential of AI in supporting pharmacological therapy.<sup>245</sup>

In addition to drug discovery, AI has shown emerging potential in the optimization of rehabilitation strategies for OP. By integrating patient-specific factors such as age, BMD, fracture risk, physical function, and comorbidities, AI-driven models may assist in designing personalized exercise and rehabilitation programs aimed at improving bone strength, balance, and functional recovery. Such as Fasihi L et al trained an ML model incorporating patients' imaging features and covariate data to generate individualized exercise recommendations aimed at improving bone health and preventing fracture risk.<sup>246</sup> However, current applications in this area remain largely exploratory, and robust clinical studies focusing on AI-guided rehabilitation for OP are still limited. In the future, with continued technological advances, AI applications in OP may evolve in a manner like those used for fracture prevention, incorporating wearable devices to continuously monitor patient status. Such systems could assist with home-based exercise programs and nutritional management, thereby supporting disease management and promoting patient rehabilitation.<sup>247</sup>

AI has also demonstrated promising applications in surgical planning and intraoperative assistance across various surgical disciplines, including orthognathic surgery.<sup>248</sup> This suggesting potential future applicability in osteoporosis-related surgical management, though current evidence specifically addressing osteoporosis surgery remains limited. In terms of postoperative prognosis prediction of OP, Klemm C et al analyzed data from 350 patients and developed four ML models capable of predicting revision surgery risk following orthopedic procedures. This demonstrates the potential of AI in refining postoperative management and long-term outcome assessment.<sup>227</sup> Nevertheless, compared with diagnostic and pharmacological applications, AI-assisted surgical approaches in OP are underrepresented in the literature, highlighting a significant gap for future research.

While current AI-driven therapeutic research in OP predominantly focuses on drug discovery, its extension to rehabilitation optimization and surgical decision support represents a promising yet underdeveloped direction. Addressing these gaps will be essential for establishing a comprehensive AI-assisted management framework that spans diagnosis, treatment, and long-term functional recovery. Recent advances indicate that AI is increasingly involved in the translation of biomaterials.<sup>249, 250</sup> Moreover, biomaterials represented by metal-organic framework nanomedicine have demonstrated significant potential in OP therapy.<sup>251</sup> Therefore, the application of AI in biomaterials translation for OP is expected to become an emerging and promising research direction. In addition, nanomaterials are currently being used as contrast agents in multimodal imaging. Their integration with AI enables personalized imaging and image-guided therapy, a strategy that may also be extended to the diagnosis and treatment of osteoporosis in the future.<sup>252</sup> The rise of LLMs has provided a more convenient approach for clinical support in OP patients, and this direction may potentially become an integral part of comprehensive management for these patients in the future.<sup>253</sup>

## Limitations

Despite these promising developments, several important limitations should be acknowledged. Considerable heterogeneity exists among published studies regarding patient populations,<sup>25,169</sup> AI algorithms,<sup>26</sup> and evaluation metrics,<sup>27</sup> which limits direct comparison and interpretation of reported performance outcomes. In addition, consistent external validation remains insufficient in many studies, raising concerns about model generalizability, robustness, and real-world clinical applicability.<sup>169,27,254</sup> Challenges related to data imbalance, and potential selection bias further highlight the need for standardized reporting frameworks and multicenter collaborative research to promote safe clinical adoption of AI technologies in osteoporosis management.<sup>25,255</sup>

## Summary

The application of AI in OP research has emerged as a rapidly evolving and interdisciplinary field. This review provides a comprehensive overview of AI technologies currently utilized in OP, categorizing existing studies into four main areas: diagnosis, genomic and bioinformatics research, risk prediction and prognosis assessment, and treatment. By understanding the current progress across these domains, clinicians and researchers can better integrate AI to achieve more intelligent, efficient, and precise management of OP.

Future research should prioritize the development of multimodal AI models that integrate imaging features with genomic and molecular data, which may improve disease characterization and address limitations in the sensitivity of certain imaging modalities, such as MRI. In addition, the use of large-scale, real-world imaging datasets for opportunistic screening represents a promising avenue for early OP detection. Emerging evidence also suggests that AI may facilitate the translation of advanced biomaterials, including metal–organic framework–based nanomedicine, enable personalized multimodal imaging and image-guided therapy through integration with nanomaterial-based contrast platforms, as well as provide comprehensive clinical support thereby expanding future therapeutic strategies for OP. Future investigations may further extend AI applications from diagnosis and drug discovery to personalized treatment planning, rehabilitation optimization, and precision therapeutic intervention. Despite these promising developments, several important limitations remain. Considerable heterogeneity exists among published studies which limit direct comparison and interpretation of reported outcomes. Moreover, insufficient external validation continues to raise concerns regarding model generalizability. Challenges related to data imbalance and potential selection bias further highlight the need for standardized reporting frameworks and multicenter collaborative research.

In conclusion, while AI has already transformed many aspects of OP research—from automated imaging analysis to biomarker discovery—its full potential in treatment personalization, biomaterials translation, and multimodal integration remains to be fully realized. Continued advances in data integration, algorithm transparency, rigorous clinical validation, and interdisciplinary collaboration will be essential for harnessing the full power of AI in the prevention and management of OP.

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## Author Contributions

All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis and interpretation, or in all these areas; took part in drafting, revising or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work.

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## References

1. Zhang J-Y, Zhong Y-H, Chen L-M, et al. Recent advance of small-molecule drugs for clinical treatment of osteoporosis: a review. *Eur J Med Chem.* **2023**;259:115654. doi:10.1016/j.ejmech.2023.115654
2. Zhu W, Guo Y, Shuai J, et al. Osteoporotic fractures prediction in Chinese postmenopausal women: a machine learning-based multi-dimensional approach. *Front Endocrinol.* **2026**;17:1788788. doi:10.3389/fendo.2026.1788788
3. Nguyen A, Lee P, Rodriguez EK, et al. Addressing the growing burden of musculoskeletal diseases in the ageing US population: challenges and innovations. *Lancet Healthy Longev.* **2025**;6(5):100707. doi:10.1016/j.lanhl.2025.100707
4. Morin SN, Leslie WD, Schousboe JT. Osteoporosis: a review. *JAMA.* **2025**;334(10):894–907. doi:10.1001/jama.2025.6003
5. About Osteoporosis | International Osteoporosis Foundation. Available from: <https://www.osteoporosis.foundation/patients/about-osteoporosis>. Accessed December 11, 2025
6. Thuillier E, Carey J, Dempsey M, et al. Virtual rehabilitation for patients with osteoporosis or other musculoskeletal disorders: a systematic review. *Virtual Real.* **2024**;28(2):93. doi:10.1007/s10055-024-00980-7
7. Lorente-Ramos R, Azpeitia-Armán J, Muñoz-Hernández A, et al. Dual-Energy X-Ray absorptiometry in the diagnosis of osteoporosis: a practical guide. *Am J Roentgenol.* **2011**;196(4):897–904. doi:10.2214/AJR.10.5416
8. Ye C, Ebeling P, Kline G. Osteoporosis. *Lancet Lond Engl.* **2025**;406(10514):2003–2016.
9. Nguyen HG, Nguyen D-T, Tran TS, et al. Artificial intelligence system for predicting areal bone mineral density from plain X-rays. *Osteoporos Int J Establ Result Coop Eur Found Osteoporos Natl Osteoporos Found USA.* **2025**;36(11):2167–2176. doi:10.1007/s00198-025-07634-7
10. Xu J, Wen X, Sun L, et al. Large model era: deep learning in osteoporosis drug discovery. *J Chem Inf Model.* **2025**;65(5):2232–2244. doi:10.1021/acs.jcim.4c02264
11. Strickland E. The turbulent past and uncertain future of AI: is there a way out of AI's boom-and-bust cycle? *IEEE Spectr.* **2021**;58(10):26–31. doi:10.1109/MSPEC.2021.9563956
12. Mathew D, Shukla VK, Chaubey A, et al. Artificial intelligence: hope for future or hype by intellectuals? In: 2021 9th Int Conf Reliab Infocom Technol Optim Trends Future Dir ICRITO. **2021**. p. 1–6
13. Li X, Zheng J, Liu C, et al. Multidimensional technological advances in cervical cancer screening: from standardized processes to precision medicine. *Biochim Biophys Acta Rev Cancer.* **2025**;1880(5):189432. doi:10.1016/j.bbcan.2025.189432
14. Liu Y, Chen S, Xiong X, et al. Artificial intelligence guided Raman spectroscopy in biomedicine: applications and prospects. *J Pharm Anal.* **2025**;15(11):101271. doi:10.1016/j.jpha.2025.101271
15. Cabello JB, Garcia VR, Torralba M, et al. Critical appraisal tools for evaluating artificial intelligence in clinical studies: scoping review. *J Med Internet Res.* **2025**;27(1):e77110. doi:10.2196/77110
16. Ekambaram S, Dokholyan NV. Peptide-based drug design using generative AI. *Chem Commun Camb Engl.* **2025**. doi:10.1039/d5cc04998a
17. Gao L, Song S, Lin J, et al. AI-Assisted design of advanced polymeric materials: challenges and solutions. *Adv Mater Deerfield Beach Fla.* **2025**;e16857.
18. Wang X, Tan T, Gao Y, et al. Mammo-AGE: deep learning estimation of breast age from mammograms. *Nat Commun.* **2025**;16(1):10934. doi:10.1038/s41467-025-65923-5
19. Barreto LS da C, Moreira Das Neves B, Bianchi J, et al. A semi-automated assessment tool for craniofacial landmarks in CBCT: inVivo7 software. *J Dent.* **2025**;165:106292. doi:10.1016/j.jdent.2025.106292
20. Abedeen A, Smith R, Hirschmann MT, et al. Synthetic computed tomography from magnetic resonance imaging: an editorial on deep learning approaches for Hip and knee image translation. *Knee Surg Sports Traumatol Arthrosc Off J ESSKA.* **2025**. doi:10.1002/ksa.70229
21. Sun Z, Li H, Zheng Y, et al. Interpretable machine learning model based on multimodal ultrasound for bedside diagnosis of acute exacerbations in COPD. *Respir Res.* **2025**;26(1):338. doi:10.1186/s12931-025-03411-6
22. Xiao H, Liang X, Li H, et al. Trends in the prevalence of osteoporosis and effects of heavy metal exposure using interpretable machine learning. *Ecotoxicol Environ Saf.* **2024**;286:117238. doi:10.1016/j.ecoenv.2024.117238
23. Zhou X, Zhao L, Zhang Z, et al. Identification of shared gene signatures and pathways for diagnosing osteoporosis with sarcopenia through integrated bioinformatics analysis and machine learning. *BMC Musculoskelet Disord.* **2024**;25(1):435. doi:10.1186/s12891-024-07555-2
24. Cho SW, Baek S, Han S, et al. Metabolic phenotyping with computed tomography deep learning for metabolic syndrome, osteoporosis and sarcopenia predicts mortality in adults. *J Cachexia Sarcopenia Muscle.* **2024**;15(4):1418–1429. doi:10.1002/jcsm.13487
25. Katagiri H, Koyano G, Katayanagi J, et al. Deep-learning framework for osteoporosis screening on low-dose X-rays: addressing image quality variability and cross-ethnic database Heterogeneity. *Eur J Radiol.* **2026**;200:112880. doi:10.1016/j.ejrad.2026.112880
26. Podoia V, Caliva F, Kazakia G, et al. Augmenting osteoporosis imaging with machine learning. *Curr Osteoporos Rep.* **2021**;19(6):699–709. doi:10.1007/s11914-021-00701-y
27. Conforti A, Ruggiero M, Lucchetti L, et al. Artificial intelligence and machine learning in the diagnosis and management of osteoporosis: a comprehensive review. *Medicina.* **2025**;62(1):27. doi:10.3390/medicina62010027
28. The evolution and impact of artificial intelligence its applications, challenges, and future directions: a review. In: IEEE Conference Publication; IEEE Xplore.
29. Das S, Roy R, Sarkar A, et al. Revolutionizing healthcare: synergistic integration of genomics, clinical decision support systems, and advanced machine learning for precision medicine. In: 2025 AI-Driven smart health soc 50. **2025**. p. 1–6.
30. Liang C, Xin S. Research status and prospects of deep learning in medical images. In: 2020 Int Conf Commun Inf Syst Comput Eng CISCE. **2020**. p. 380–382.
31. Clive C, Singh A, Overmeer B, et al. Large-Scale automated phenotyping of cardiac arrest and withdrawal of life-sustaining therapy using electronic health record data. *Resuscitation.* **2025**;218:110919. doi:10.1016/j.resuscitation.2025.110919

32. Nghiem E, Ziguoras S, Yu H, et al. Combined endoscopic and robotic-assisted transcolonic polypectomy as a novel technique for challenging polyp resection. *Surg Endosc*. 2025;40(1):810–814. doi:10.1007/s00464-025-12445-2
33. Riener R, Nef T, Colombo G. Robot-aided neurorehabilitation of the upper extremities. *Med Biol Eng Comput*. 2005;43(1):2–10. doi:10.1007/BF02345116
34. LeCun Y, Bengio Y, Hinton G. Deep learning. *Nature*. 2015;521(7553):436–444. doi:10.1038/nature14539
35. Tian C, Cheng T, Peng Z, et al. A survey on deep learning fundamentals. *Artif Intell Rev*. 2025;58(12):381.
36. Dong S, Wang P, Abbas K. A survey on deep learning and its applications. *Comput Sci Rev*. 2021;40:100379. doi:10.1016/j.cosrev.2021.100379
37. Murphy KP. *Machine Learning: A Probabilistic Perspective*. The MIT Press; 2012.
38. Domingos P. A few useful things to know about machine learning. *Commun ACM*. 2012;55(10):78–87. doi:10.1145/2347736.2347755
39. What is Machine Learning? | IBM. 2025. Available from: <https://www.ibm.com/think/topics/machine-learning>. Accessed December 14, 2025.
40. Nian S, Zhao Y, Li C, et al. Development and validation of a radiomics-based model for predicting osteoporosis in patients with lumbar compression fractures. *Spine J Off J North Am Spine Soc*. 2024;24(9):1625–1634. doi:10.1016/j.spinee.2024.04.016
41. Oh J, Kim B, Oh G, et al. End-to-End semi-supervised opportunistic osteoporosis screening using computed tomography. *Endocrinol Metab Seoul Korea*. 2024;39(3):500–510. doi:10.3803/EnM.2023.1860
42. Hung TNK, Le NQK, Le NH, et al. An AI-based prediction model for drug-drug interactions in osteoporosis and paget's diseases from SMILES. *Mol Inform*. 2022;41(6):e2100264. doi:10.1002/minf.202100264
43. Zheng Z, Zhang X, Oh B-K, et al. Identification of combined biomarkers for predicting the risk of osteoporosis using machine learning. *Aging*. 2022;14(10):4270–4280. doi:10.18632/aging.204084
44. Shim J-G, Kim DW, Ryu K-H, et al. Application of machine learning approaches for osteoporosis risk prediction in postmenopausal women. *Arch Osteoporos*. 2020;15(1):169. doi:10.1007/s11657-020-00802-8
45. Tarighatnia A, Amanzadeh M, Hamedan M, et al. Deep learning-based evaluation of panoramic radiographs for osteoporosis screening: a systematic review and meta-analysis. *BMC Med Imaging*. 2025;25(1):86. doi:10.1186/s12880-025-01626-z
46. Paderno A, Ataide Gomes EJ, Gilberg L, et al. Artificial intelligence-enhanced opportunistic screening of osteoporosis in CT scan: a scoping review. *Osteoporos Int J Establ Result Coop Eur Found Osteoporos Natl Osteoporos Found USA*. 2024;35(10):1681–1692. doi:10.1007/s00198-024-07179-1
47. Kavitha MS, An S-Y, An C-H, et al. Texture analysis of mandibular cortical bone on digital dental panoramic radiographs for the diagnosis of osteoporosis in Korean women. *Oral Surg Oral Med Oral Pathol Oral Radiol*. 2015;119(3):346–356. doi:10.1016/j.oooo.2014.11.009
48. Gao L, Jiao T, Feng Q, et al. Application of artificial intelligence in diagnosis of osteoporosis using medical images: a systematic review and meta-analysis. *Osteoporos Int J Establ Result Coop Eur Found Osteoporos Natl Osteoporos Found USA*. 2021;32(7):1279–1286. doi:10.1007/s00198-021-05887-6
49. Galbusera F, Cina A, O'Riordan D, et al. Estimating lumbar bone mineral density from conventional MRI and radiographs with deep learning in spine patients. *Eur Spine J Off Publ Eur Spine Soc Eur Spinal Deform Soc Eur Sect Cerv Spine Res Soc*. 2024;33(11):4092–4103. doi:10.1007/s00586-024-08463-8
50. Hu M, Zou L, Lu J, et al. Construction of a 5-feature gene model by support vector machine for classifying osteoporosis samples. *Bioengineered*. 2021;12(1):6821–6830. doi:10.1080/21655979.2021.1971026
51. Zhang X, Zhu X, Gu W, et al. Elucidating the mechanism of phthalates induced osteoporosis through network toxicology and molecular docking. *Ecotoxicol Environ Saf*. 2025;291:117820. doi:10.1016/j.ecoenv.2025.117820
52. Yang J, Yang W, Hu Y, et al. Screening of genes co-associated with osteoporosis and chronic HBV infection based on bioinformatics analysis and machine learning. *Front Immunol*. 2024;15:1472354. doi:10.3389/fimmu.2024.1472354
53. Chang C-Y, Peng C-H, Chen F-Y, et al. The risk factors determined by four machine learning methods for the change of difference of bone mineral density in post-menopausal women after three years follow-up. *Sci Rep*. 2024;14(1):23234. doi:10.1038/s41598-024-73799-6
54. Xu R, Chen Y, Yao Z, et al. Application of machine learning algorithms to identify people with low bone density. *Front Public Health*. 2024;12:1347219. doi:10.3389/fpubh.2024.1347219
55. Yang X, Du Z, Xing S. CHST3, PGBD5, and SLIT2 can be identified as potential genes for the diagnosis and treatment of osteoporosis and sarcopenia. *Sci Rep*. 2025;15(1):374. doi:10.1038/s41598-024-83231-8
56. Li J, Zhang P, Xu J, et al. Prediction of bone mineral density based on computer tomography images using deep learning model. *Gerontology*. 2025;71(1):71–80. doi:10.1159/000542396
57. Ho C-S, Fan T-Y, Kuo C-F, et al. HardNet-based deep learning model for osteoporosis screening and bone mineral density inference from hand radiographs. *Bone*. 2025;190:117317. doi:10.1016/j.bone.2024.117317
58. Pan J, Lin P-C, Gong S-C, et al. Feasibility study of opportunistic osteoporosis screening on chest CT using a multi-feature fusion DCNN model. *Arch Osteoporos*. 2024;19(1):98. doi:10.1007/s11657-024-01455-7
59. Woo BFY, Cato K, Cho H, et al. The use of large language models in clinical documentation: a scoping review. *Int J Nurs Stud*. 2025;176:105322. doi:10.1016/j.ijnurstu.2025.105322
60. Kong SH. Incorporating artificial intelligence into fracture risk assessment: using clinical imaging to predict the unpredictable. *Endocrinol Metab Seoul Korea*. 2025;40(4):499–507. doi:10.3803/EnM.2025.2518
61. Li D, Liu F, Hou Y, et al. Changes in periprosthetic bone mineral density following arthroplasty: an in-depth review and current perspectives. *Curr Osteoporos Rep*. 2025;23(1):30. doi:10.1007/s11914-025-00921-6
62. Johnson PM, Umapathy L, Gigax B, et al. Artificial intelligence in prostate MRI: addressing current limitations through emerging technologies. *J Magn Reson Imaging JMRI*. 2025. doi:10.1002/jmri.70189
63. Zhang W, Tai D, Kang S, et al. The application progress of artificial intelligence in osteoporosis diagnosis. *Front Artif Intell*. 2025;8:1699762. doi:10.3389/frai.2025.1699762
64. Pulkkinen P, Saarakkala S, Nieminen MT, et al. Standard radiography: untapped potential in the assessment of osteoporotic fracture risk. *Eur Radiol*. 2013;23(5):1375–1382. doi:10.1007/s00330-012-2722-9
65. Xue L, Qin G, Chang S, et al. Osteoporosis prediction in lumbar spine X-ray images using the multi-scale weighted fusion contextual transformer network. *Artif Intell Med*. 2023;143:102639. doi:10.1016/j.artmed.2023.102639

66. Jang M, Kim M, Bae SJ, et al. Opportunistic osteoporosis screening using chest radiographs with deep learning: development and external validation with a cohort dataset. *J Bone Miner Res off J Am Soc Bone Miner Res.* **2022**;37(2):369–377. doi:10.1002/jbmr.4477
67. Zhang B, Yu K, Ning Z, et al. Deep learning of lumbar spine X-ray for osteopenia and osteoporosis screening: a multicenter retrospective cohort study. *Bone.* **2020**;140:115561. doi:10.1016/j.bone.2020.115561
68. Saphagirivasan V, Anburajan M. Diagnosis of osteoporosis by extraction of trabecular features from Hip radiographs using support vector machine: an investigation panorama with DXA. *Comput Biol Med.* **2013**;43(11):1910–1919. doi:10.1016/j.combiomed.2013.09.002
69. Areecal AS, Jayasheelan N, Kamath J, et al. Early diagnosis of osteoporosis using radiogrammetry and texture analysis from hand and wrist radiographs in Indian population. *Osteoporos Int J Establ Result Coop Eur Found Osteoporos Natl Osteoporos Found USA.* **2018**;29(3):665–673. doi:10.1007/s00198-017-4328-1
70. Gharehmohammadi F, Sebros R. Deep learning opportunistic screening for osteoporosis and osteopenia using radiographs of the foot or ankle - A pilot study. *Eur J Radiol.* **2025**;184:111980. doi:10.1016/j.ejrad.2025.111980
71. Naguib SM, Saleh MK, Hamza HM, et al. A new superfluity deep learning model for detecting knee osteoporosis and osteopenia in X-ray images. *Sci Rep.* **2024**;14(1):25434. doi:10.1038/s41598-024-75549-0
72. Xie H, Gu C, Zhang W, et al. A few-shot learning framework for the diagnosis of osteopenia and osteoporosis using knee X-ray images. *J Int Med Res.* **2024**;52(9):3000605241274576. doi:10.1177/03000605241274576
73. Sarmadi A, Razavi ZS, van Wijnen AJ, et al. Comparative analysis of vision transformers and convolutional neural networks in osteoporosis detection from X-ray images. *Sci Rep.* **2024**;14(1):18007. doi:10.1038/s41598-024-69119-7
74. Srinivasan D, Kiran A, Parameswari S, et al. Bonevoyage: navigating the depths of osteoporosis detection with a dual-core ensemble of cascaded ShuffleNet and neural networks. *J X-Ray Sci Technol.* **2025**;33(1):3–25.
75. Muhammad I, Rao RS, Lee B. BONE-Net: a novel hybrid deep-learning model for effective osteoporosis detection. *PLoS One.* **2025**;20(10):e0334664. doi:10.1371/journal.pone.0334664
76. Qureshi MB, Sani M, Raza A, et al. Deep-learning based osteoporosis classification in knee X-rays using transfer-learning approach. *Sci Rep.* **2025**;15(1):38448. doi:10.1038/s41598-025-24338-4
77. Lin C, Tsai D-J, Wang -C-C, et al. Osteoporotic precise screening using chest radiography and artificial neural network: the OPSCAN randomized controlled trial. *Radiology.* **2024**;311(3):e231937. doi:10.1148/radiol.231937
78. Asamoto T, Takegami Y, Sato Y, et al. External validation of a deep learning model for predicting bone mineral density on chest radiographs. *Arch Osteoporos.* **2024**;19(1):15. doi:10.1007/s11657-024-01372-9
79. Tsai D-J, Lin C, Lin C-S, et al. Artificial intelligence-enabled chest X-ray classifies osteoporosis and identifies mortality risk. *J Med Syst.* **2024**;48(1):12. doi:10.1007/s10916-023-02030-2
80. Ohta Y, Yamamoto K, Katayama Y, et al. Development of AI model for dual detection of low bone mineral density in the femoral neck and lumbar vertebrae using chest radiographs. *J Clin Densitom off J Int Soc Clin Densitom.* **2025**;28(4):101604. doi:10.1016/j.jocd.2025.101604
81. Tang J, Yin X, Lai J, et al. Automatic opportunistic osteoporosis screening using chest X-ray images via deep neural networks. *Bone.* **2025**;201:117618. doi:10.1016/j.bone.2025.117618
82. Iwao Y, Shiotsuki K, Hashimoto F, et al. An exploratory study of explainable deep learning for predicting bone mineral density using clavicle features on chest radiographs: a multi-task approach with regression and segmentation. *J Appl Clin Med Phys.* **2025**;26(12):e70336. doi:10.1002/acm2.70336
83. Park J, Kim N-Y, Bae H-J, et al. Opportunistic screening of low bone mass using knowledge distillation-based deep learning in chest X-rays with external validations. *Arch Osteoporos.* **2025**;20(1):131. doi:10.1007/s11657-025-01609-1
84. Lee S, Choe EK, Kang HY, et al. The exploration of feature extraction and machine learning for predicting bone density from simple spine X-ray images in a Korean population. *Skeletal Radiol.* **2020**;49(4):613–618. doi:10.1007/s00256-019-03342-6
85. Hsieh C-I, Zheng K, Lin C, et al. Automated bone mineral density prediction and fracture risk assessment using plain radiographs via deep learning. *Nat Commun.* **2021**;12(1):5472. doi:10.1038/s41467-021-25779-x
86. Mao L, Xia Z, Pan L, et al. Deep learning for screening primary osteopenia and osteoporosis using spine radiographs and patient clinical covariates in a Chinese population. *Front Endocrinol.* **2022**;13:971877. doi:10.3389/fendo.2022.971877
87. Dong Q, Luo G, Lane NE, et al. Deep learning classification of spinal osteoporotic compression fractures on radiographs using an adaptation of the genant semiquantitative criteria. *Acad Radiol.* **2022**;29(12):1819–1832. doi:10.1016/j.acra.2022.02.020
88. Hong N, Cho SW, Shin S, et al. Deep-Learning-Based detection of vertebral fracture and osteoporosis using lateral spine X-Ray radiography. *J Bone Miner Res Off J Am Soc Bone Miner Res.* **2023**;38(6):887–895. doi:10.1002/jbmr.4814
89. Dong Q, Luo G, Lane NE, et al. Generalizability of deep learning classification of spinal osteoporotic compression fractures on radiographs using an adaptation of the modified-2 algorithm-based qualitative criteria. *Acad Radiol.* **2023**;30(12):2973–2987. doi:10.1016/j.acra.2023.04.023
90. Moro T, Yoshimura N, Saito T, et al. Development of artificial intelligence-assisted lumbar and femoral BMD estimation system using anteroposterior lumbar X-Ray images. *J Orthop Res Off Publ Orthop Res Soc.* **2025**;43(9):1619–1631. doi:10.1002/jor.70000
91. Hong N, Cho SW, Lee YH, et al. Deep learning-based identification of vertebral fracture and osteoporosis in lateral spine radiographs and DXA vertebral fracture assessment to predict incident fracture. *J Bone Miner Res Off J Am Soc Bone Miner Res.* **2025**;40(5):628–638.
92. Hong J, Sung H, Choi J-O, et al. A prediction model of pediatric bone density from plain spine radiographs using deep learning. *Sci Rep.* **2025**;15(1):13039.
93. Tamai K, Imanishi K, Terakawa M, et al. Deep learning algorithm for identifying osteopenia/osteoporosis using cervical radiography. *Sci Rep.* **2025**;15(1):25274. doi:10.1038/s41598-025-11285-3
94. Boonrod A, Kittipongphat N, Piyaapapaphan P, et al. Assessing deep learning model performance in osteoporosis screening with lumbar spine radiographs. *J Bone Miner Metab.* **2026**;44(1):124–131. doi:10.1007/s00774-025-01672-1
95. Kavitha MS, Ganesh Kumar P, Park S-Y, et al. Automatic detection of osteoporosis based on hybrid genetic swarm fuzzy classifier approaches. *Dento Maxillo Facial Radiol.* **2016**;45(7):20160076. doi:10.1259/dmfr.20160076
96. Hwang JJ, Lee J-H, Han -S-S, et al. Strut analysis for osteoporosis detection model using dental panoramic radiography. *Dento Maxillo Facial Radiol.* **2017**;46(7):20170006. doi:10.1259/dmfr.20170006

97. Lee J-S, Adhikari S, Liu L, et al. Osteoporosis detection in panoramic radiographs using a deep convolutional neural network-based computer-assisted diagnosis system: a preliminary study. *Dento Maxillo Facial Radiol.* **2019**;48(1):20170344. doi:10.1259/dmfr.20170344
98. Alzubaidi MA, Ootom M. A comprehensive study on feature types for osteoporosis classification in dental panoramic radiographs. *Comput Methods Programs Biomed.* **2020**;188:105301.
99. Nakamoto T, Taguchi A, Kakimoto N. Osteoporosis screening support system from panoramic radiographs using deep learning by convolutional neural network. *Dento Maxillo Facial Radiol.* **2022**;51(6):20220135.
100. Tassoker M, Öziç MÜ, Yuce F. Comparison of five convolutional neural networks for predicting osteoporosis based on mandibular cortical index on panoramic radiographs. *Dento Maxillo Facial Radiol.* **2022**;51(6):20220108. doi:10.1259/dmfr.20220108
101. Sukegawa S, Fujimura A, Taguchi A, et al. Identification of osteoporosis using ensemble deep learning model with panoramic radiographs and clinical covariates. *Sci Rep.* **2022**;12(1):6088. doi:10.1038/s41598-022-10150-x
102. Yadalam PK, Pawar AR, Natarajan PM, et al. A novel dual embedding few-shot learning approach for classifying bone loss using orthopantomogram radiographic notes. *Head Face Med.* **2025**;21(1):49. doi:10.1186/s13005-025-00528-3
103. Roebbers P, Schulze R. Use of neural networks to predict a potential osteoporotic metabolic condition based on the panoramic mandibular index. *Oral Surg Oral Med Oral Pathol Oral Radiol.* **2026**;141(2):245–254. doi:10.1016/j.oooo.2025.09.022
104. Yamamoto N, Sukegawa S, Kitamura A, et al. Deep learning for osteoporosis classification using hip radiographs and patient clinical covariates. *Biomolecules.* **2020**;10(11):1534. doi:10.3390/biom10111534
105. Jang R, Choi JH, Kim N, et al. Prediction of osteoporosis from simple Hip radiography using deep learning algorithm. *Sci Rep.* **2021**;11(1):19997. doi:10.1038/s41598-021-99549-6
106. Yamamoto N, Sukegawa S, Yamashita K, et al. Effect of patient clinical variables in osteoporosis classification using hip X-rays in deep learning analysis. *Med Kaunas Lith.* **2021**;57(8):846.
107. Nguyen TP, Chae D-S, Park S-J, et al. A novel approach for evaluating bone mineral density of hips based on Sobel gradient-based map of radiographs utilizing convolutional neural network. *Comput Biol Med.* **2021**;132:104298. doi:10.1016/j.combiomed.2021.104298
108. Acosta-Batlle J, Coronado-Gutiérrez D, Soto J, et al. An AI-based pipeline for osteoporosis/osteopenia prediction using hip radiographs. *Skeletal Radiol.* **2026**;55(2):415–424. doi:10.1007/s00256-025-05033-x
109. Singh A, Dutta MK, Jennane R, et al. Classification of the trabecular bone structure of osteoporotic patients using machine vision. *Comput Biol Med.* **2017**;91:148–158.
110. Osteoporosis prediction from hand X-ray images using segmentation-for-classification and self-supervised learning. Available from: <https://pubmed.ncbi.nlm.nih.gov/41006333/>. Accessed May 9, 2026.
111. Lee K-XA, Chang H-C, Chiu Y-C, et al. Enhanced osteoporosis screening via multi-output deep learning: segmentation and classification of metacarpal radiographs. *Eur J Radiol.* **2026**;194:112507. doi:10.1016/j.ejrad.2025.112507
112. Liu J, Wang J, Ruan W, et al. Diagnostic and gradation model of osteoporosis based on improved deep U-net network. *J Med Syst.* **2019**;44(1):15. doi:10.1007/s10916-019-1502-3
113. Zhou K, Zhu Y, Luo X, et al. A novel hybrid deep learning framework based on biplanar X-ray radiography images for bone density prediction and classification. *Osteoporos Int J Establ Result Coop Eur Found Osteoporos Natl Osteoporos Found USA.* **2025**;36(3):521–530. doi:10.1007/s00198-024-07378-w
114. Yen T-Y, Ho C-S, Pei Y-C, et al. Predicting osteoporosis from kidney-ureter-bladder radiographs utilizing deep convolutional neural networks. *Bone.* **2024**;184:117107. doi:10.1016/j.bone.2024.117107
115. Quagliato L, Seo J, Hong J, et al. Synthetic data-enhanced classification of prevalent osteoporotic fractures using dual-energy X-Ray absorptiometry-based geometric and material parameters. *Endocrinol Metab Seoul Korea.* **2025**;40(3):484–497. doi:10.3803/EnM.2024.2211
116. Chen RJ, Wang JJ, Williamson DFK, et al. Algorithmic fairness in artificial intelligence for medicine and healthcare. *Nat Biomed Eng.* **2023**;7(6):719–742.
117. Zhao Y, Qiu J, Li Y, et al. Machine-learning models for diagnosis of rotator cuff tears in osteoporosis patients based on anteroposterior X-rays of the shoulder joint. *SLAS Technol.* **2024**;29(4):100149. doi:10.1016/j.slast.2024.100149
118. Ashok Kumar D, Anburajan M, Snehalatha U. Evaluation of low bone mass and prediction of fracture risk using metacarpal radiogrammetry method: a comparative study with DXA and X-ray phantom. *Int J Rheum Dis.* **2018**;21(7):1350–1371. doi:10.1111/1756-185X.13326
119. Troy KL, Edwards WB. Practical considerations for obtaining high quality quantitative computed tomography data of the skeletal system. *Bone.* **2018**;110:58–65. doi:10.1016/j.bone.2018.01.013
120. Tomita N, Cheung YY, Hassanpour S. Deep neural networks for automatic detection of osteoporotic vertebral fractures on CT scans. *Comput Biol Med.* **2018**;98:8–15. doi:10.1016/j.combiomed.2018.05.011
121. Lim HK, Ha HI, Park S-Y, et al. Prediction of femoral osteoporosis using machine-learning analysis with radiomics features and abdomen-pelvic CT: a retrospective single center preliminary study. *PLoS One.* **2021**;16(3):e0247330. doi:10.1371/journal.pone.0247330
122. Yasaka K, Akai H, Kunitatsu A, et al. Prediction of bone mineral density from computed tomography: application of deep learning with a convolutional neural network. *Eur Radiol.* **2020**;30(6):3549–3557. doi:10.1007/s00330-020-06677-0
123. Namatevs I, Nikulins A, Edelmers E, et al. Modular neural networks for osteoporosis detection in mandibular cone-beam computed tomography scans. *Tomogr Ann Arbor Mich.* **2023**;9(5):1772–1786.
124. Zhang K, Lin P-C, Pan J, et al. DeepmdQCT: a multitask network with domain invariant features and comprehensive attention mechanism for quantitative computer tomography diagnosis of osteoporosis. *Comput Biol Med.* **2024**;170:107916.
125. Yang J, Liao M, Wang Y, et al. Opportunistic osteoporosis screening using chest CT with artificial intelligence. *Osteoporos Int J Establ Result Coop Eur Found Osteoporos Natl Osteoporos Found USA.* **2022**;33(12):2547–2561. doi:10.1007/s00198-022-06491-y
126. Breit H-C, Varga-Szemes A, Schoepf UJ, et al. CNN-based evaluation of bone density improves diagnostic performance to detect osteopenia and osteoporosis in patients with non-contrast chest CT examinations. *Eur J Radiol.* **2023**;161:110728. doi:10.1016/j.ejrad.2023.110728
127. Peng T, Zeng X, Li Y, et al. A study on whether deep learning models based on CT images for bone density classification and prediction can be used for opportunistic osteoporosis screening. *Osteoporos Int J Establ Result Coop Eur Found Osteoporos Natl Osteoporos Found USA.* **2024**;35(1):117–128. doi:10.1007/s00198-023-06900-w
128. Park H, Kang WY, Woo OH, et al. Automated deep learning-based bone mineral density assessment for opportunistic osteoporosis screening using various CT protocols with multi-vendor scanners. *Sci Rep.* **2024**;14(1):25014. doi:10.1038/s41598-024-73709-w

129. Lin X, Shen R, Zheng X, et al. Utilizing radiomics techniques to isolate a single vertebral body from chest CT for opportunistic osteoporosis screening. *BMC Musculoskelet Disord.* 2024;25(1):785. doi:10.1186/s12891-024-07903-2
130. Fang K, Zheng X, Lin X, et al. A comprehensive approach for osteoporosis detection through chest CT analysis and bone turnover markers: harnessing radiomics and deep learning techniques. *Front Endocrinol.* 2024;15:1296047. doi:10.3389/fendo.2024.1296047
131. Tong X, Wang S, Cheng Q, et al. Effect of fully automatic classification model from different tube voltage images on bone density screening: a self-controlled study. *Eur J Radiol.* 2024;177:111521. doi:10.1016/j.ejrad.2024.111521
132. Wang S, Tong X, Fan Y, et al. Combining deep learning and radiomics for automated, objective, comprehensive bone mineral density assessment from low-dose chest computed tomography. *Acad Radiol.* 2024;31(3):1180–1188. doi:10.1016/j.acra.2023.08.030
133. Li Y, Liu S, Zhang Y, et al. Deep learning-enhanced opportunistic osteoporosis screening in ultralow-voltage (80 kV) chest CT: a preliminary study. *Acad Radiol.* 2025;32(7):4254–4265. doi:10.1016/j.acra.2024.11.062
134. Li Y, Zhu Y, Zhang Y, et al. Deep learning-based automated opportunistic osteoporosis screening using chest LDCT and lumbar CT: a Multicenter Cohort Study. *Acad Radiol.* 2025;32(12):7382–7396. doi:10.1016/j.acra.2025.09.015
135. Zhou K, Xin E, Yang S, et al. Automated fast prediction of bone mineral density from low-dose computed tomography. *Acad Radiol.* 2025;32(7):4111–4120. doi:10.1016/j.acra.2025.02.041
136. Kuo D-P, Chen Y-C, Cheng S-J, et al. A vision transformer-convolutional neural network framework for decision-transparent dual-energy X-ray absorptiometry recommendations using chest low-dose CT. *Int J Med Inf.* 2025;199:105901. doi:10.1016/j.ijmedinf.2025.105901
137. Li Y, Ye K, Liu S, et al. Deep learning-enhanced opportunistic osteoporosis screening in 100 kV low-voltage chest CT: a novel way toward bone mineral density measurement and radiation dose reduction. *Acad Radiol.* 2025;32(11):6812–6822. doi:10.1016/j.acra.2025.07.060
138. Wei L, Qiu Y, Lin W, et al. Combination of artificial intelligence and chest computed tomography to assess bone mineral density. *Skeletal Radiol.* 2026;55(3):671–680. doi:10.1007/s00256-025-05067-1
139. Zhang Y, Shao R, Zhang K, et al. Deep learning model for osteoporosis screening on chest CT with low tube voltage. *Eur Spine J off Publ Eur Spine Soc Eur Spinal Deform Soc Eur Sect Cerv Spine Res Soc.* 2026;35(3):1230–1237. doi:10.1007/s00586-025-09540-2
140. Tariq A, Patel BN, Sensakovic WF, et al. Opportunistic screening for low bone density using abdominopelvic computed tomography scans. *Med Phys.* 2023;50(7):4296–4307. doi:10.1002/mp.16230
141. Yuan X, Liang Y, Yang H, et al. Applying machine learning analysis based on proximal femur of abdominal computed tomography to screen for abnormal bone mass in femur. *Acad Radiol.* 2024;31(5):2003–2010. doi:10.1016/j.acra.2023.10.035
142. Adarve-Castro A, Soria-Utrilla V, Castro-García JM, et al. Advanced radiomic prediction of osteoporosis in primary hyperparathyroidism: a machine learning-based analysis of CT images. *Radiol Med.* 2025;130(7):1084–1091. doi:10.1007/s11547-025-02004-z
143. Paukovitch M, Fechner T, Felbel D, et al. Opportunistic computed tomography (CT) assessment of osteoporosis in patients undergoing transcatheter aortic valve replacement (TAVR). *Arch Osteoporos.* 2025;20(1):100. doi:10.1007/s11657-025-01579-4
144. Sarquis Serpa A, Straus Takahashi M, Júdece de Mattos Farina EM, et al. Advancing osteoporosis opportunistic screening: multicenter validation of a deep learning algorithm using abdominal CT scans. *Abdom Radiol.* 2026;51(5):2631–2641. doi:10.1007/s00261-025-05213-2
145. Tang C, Zhang W, Li H, et al. CNN-based qualitative detection of bone mineral density via diagnostic CT slices for osteoporosis screening. *Osteoporos Int J Establ Result Coop Eur Found Osteoporos Natl Osteoporos Found USA.* 2021;32(5):971–979. doi:10.1007/s00198-020-05673-w
146. Dzierżak R, Omiotek Z. Application of deep convolutional neural networks in the diagnosis of osteoporosis. *Sensors.* 2022;22(21):8189. doi:10.3390/s22218189
147. Cheng L, Cai F, Xu M, et al. A diagnostic approach integrated multimodal radiomics with machine learning models based on lumbar spine CT and X-ray for osteoporosis. *J Bone Miner Metab.* 2023;41(6):877–889. doi:10.1007/s00774-023-01469-0
148. Wang J, He Y, Yan L, et al. Predicting osteoporosis and osteopenia by fusing deep transfer learning features and classical radiomics features based on single-source dual-energy CT imaging. *Acad Radiol.* 2024;31(10):4159–4170. doi:10.1016/j.acra.2024.04.022
149. Chen B, Cui J, Li C, et al. Application of radiomics model based on lumbar computed tomography in diagnosis of elderly osteoporosis. *J Orthop Res Off Publ Orthop Res Soc.* 2024;42(6):1356–1368. doi:10.1002/jor.25789
150. Liu J, Wang H, Shan X, et al. Hybrid transformer convolutional neural network-based radiomics models for osteoporosis screening in routine CT. *BMC Med Imaging.* 2024;24(1):62. doi:10.1186/s12880-024-01240-5
151. Küçükçiloğlu Y, Şekeroğlu B, Adalı T, et al. Prediction of osteoporosis using MRI and CT scans with unimodal and multimodal deep-learning models. *Diagn Interv Radiol Ank Turk.* 2024;30(1):9–20. doi:10.4274/dir.2023.232116
152. Xu Y, Li D, Chen Q, et al. Full supervised learning for osteoporosis diagnosis using micro-CT images. *Microsc Res Tech.* 2013;76(4):333–341. doi:10.1002/jemt.22171
153. Sebro R, De la Garza-Ramos C, Peterson JJ. Detecting whether L1 or other lumbar levels would be excluded from DXA bone mineral density analysis during opportunistic CT screening for osteoporosis using machine learning. *Int J Comput Assist Radiol Surg.* 2023;18(12):2261–2272. doi:10.1007/s11548-023-02910-5
154. Du C, He J, Cheng Q, et al. Automated opportunistic screening for osteoporosis using deep learning-based automatic segmentation and radiomics on proximal femur images from low-dose abdominal CT. *BMC Musculoskelet Disord.* 2025;26(1):378. doi:10.1186/s12891-025-08631-x
155. Pan Y, Shi D, Wang H, et al. Automatic opportunistic osteoporosis screening using low-dose chest computed tomography scans obtained for lung cancer screening. *Eur Radiol.* 2020;30(7):4107–4116. doi:10.1007/s00330-020-06679-y
156. Pan Y, Zhao F, Cheng G, et al. Automated vertebral bone mineral density measurement with phantomless internal calibration in chest LDCT scans using deep learning. *Br J Radiol.* 2023;96(1152):20230047. doi:10.1259/bjr.20230047
157. Hathaway QA, Kasaian A, Pan T, et al. A deep learning model for three-dimensional determination of whole thoracic vertebral bone mineral density from noncontrast chest CT: the Multi-Ethnic Study of Atherosclerosis. *Radiology.* 2025;314(3):e242133. doi:10.1148/radiol.242133
158. Fang Y, Li W, Chen X, et al. Opportunistic osteoporosis screening in multi-detector CT images using deep convolutional neural networks. *Eur Radiol.* 2021;31(4):1831–1842. doi:10.1007/s00330-020-07312-8
159. Oh S, Kang WY, Park H, et al. Evaluation of deep learning-based quantitative computed tomography for opportunistic osteoporosis screening. *Sci Rep.* 2024;14(1):363. doi:10.1038/s41598-023-45824-7

160. Li Y, Jin D, Zhang Y, et al. Utilizing artificial intelligence to determine bone mineral density using spectral CT. *Bone*. 2025;192:117321. doi:10.1016/j.bone.2024.117321
161. Schmidt D, Ulén J, Enqvist O, et al. Deep learning takes the pain out of back breaking work - Automatic vertebral segmentation and attenuation measurement for osteoporosis. *Clin Imaging*. 2022;81:54–59. doi:10.1016/j.clinimag.2021.08.009
162. Biamonte E, Levi R, Carrone F, et al. Artificial intelligence-based radiomics on computed tomography of lumbar spine in subjects with fragility vertebral fractures. *J Endocrinol Invest*. 2022;45(10):2007–2017. doi:10.1007/s40618-022-01837-z
163. Zhang J, Xia L, Zhang X, et al. Development and validation of a predictive model for vertebral fracture risk in osteoporosis patients. *Eur Spine J Off Publ Eur Spine Soc Eur Spinal Deform Soc Eur Sect Cerv Spine Res Soc*. 2024;33(8):3242–3260. doi:10.1007/s00586-024-08235-4
164. Yoshida K, Tanabe Y, Nishiyama H, et al. Feasibility of bone mineral density and bone microarchitecture assessment using deep learning with a convolutional neural network. *J Comput Assist Tomogr*. 2023;47(3):467–474. doi:10.1097/RCT.0000000000001437
165. Feng E, Jayasuriya NM, Nathani KR, et al. Artificial intelligence image analysis for Hounsfield units in preoperative thoracolumbar CT scans: an automated screening for osteoporosis in patients undergoing spine surgery. *J Neurosurg Spine*. 2025;43(1):1–8. doi:10.3171/2025.1.SPINE24900
166. Song J, Cho SW, Yoo HJ, et al. Enhanced opportunistic CT screening for osteoporosis using Machine learning derived volumetric vertebral and complementary body composition information. *Eur J Radiol*. 2026;194:112555. doi:10.1016/j.ejrad.2025.112555
167. Hiyama A, Sakai D, Katoh H, et al. Osteoporosis prediction using lumbar CT hounsfield units: comparative performance and clinical implications of seven machine learning models. *Eur Spine J Off Publ Eur Spine Soc Eur Spinal Deform Soc Eur Sect Cerv Spine Res Soc*. 2026;35(3):1149–1161. doi:10.1007/s00586-026-09753-z
168. Wu Y, Yang X, Wang M, et al. Artificial intelligence assisted automatic screening of opportunistic osteoporosis in computed tomography images from different scanners. *Eur Radiol*. 2025;35(4):2287–2295. doi:10.1007/s00330-024-11046-2
169. Ong W, Liu RW, Makmur A, et al. Artificial intelligence applications for osteoporosis classification using computed tomography. *Bioeng Basel Switz*. 2023;10(12):1364.
170. Liu Z, Li Y, Zhang C, et al. Radiomics and machine learning for osteoporosis detection using abdominal computed tomography: a retrospective multicenter study. *BMC Med Imaging*. 2025;25(1):235. doi:10.1186/s12880-025-01743-9
171. Park C-S, Kang S-R, Kim J-E, et al. Validation of bone mineral density measurement using quantitative CBCT image based on deep learning. *Sci Rep*. 2023;13(1):11921. doi:10.1038/s41598-023-38943-8
172. Praveen AD, Sollmann N, Baum T, et al. CT image-based biomarkers for opportunistic screening of osteoporotic fractures: a systematic review and meta-analysis. *Osteoporos Int J Establ Result Coop Eur Found Osteoporos Natl Osteoporos Found USA*. 2024;35(6):971–996. doi:10.1007/s00198-024-07029-0
173. Huppertz A, Radmer S, Asbach P, et al. Computed tomography for preoperative planning in minimal-invasive total Hip arthroplasty: radiation exposure and cost analysis. *Eur J Radiol*. 2011;78(3):406–413. doi:10.1016/j.ejrad.2009.11.024
174. Ferizi U, Besser H, Hysi P, et al. Artificial intelligence applied to osteoporosis: a performance comparison of machine learning algorithms in predicting fragility fractures from MRI data. *J Magn Reson Imaging JMRI*. 2019;49(4):1029–1038. doi:10.1002/jmri.26280
175. Vogl F, Friesenbichler B, Hüsken L, et al. Can low-frequency guided waves at the tibia paired with machine learning differentiate between healthy and osteopenic/osteoporotic subjects? A pilot study. *Ultrasonics*. 2019;94:109–116. doi:10.1016/j.ultras.2018.11.012
176. Mohanty K, Yousefian O, Karbalaiesadegh Y, et al. Artificial neural network to estimate micro-architectural properties of cortical bone using ultrasonic attenuation: a 2-D numerical study. *Comput Biol Med*. 2019;114:103457. doi:10.1016/j.compbiomed.2019.103457
177. Ferguson HE, Herickhoff CD, Viano AM, et al. Evaluating bone density with ultrasonic backscatter: leveraging time-frequency analyses and convolutional neural networks. *Ultrasonics*. 2026;159:107845. doi:10.1016/j.ultras.2025.107845
178. Thomsen K, Jepsen DB, Matzen L, et al. Is calcaneal quantitative ultrasound useful as a prescreen stratification tool for osteoporosis? *Osteoporos Int J Establ Result Coop Eur Found Osteoporos Natl Osteoporos Found USA*. 2015;26(5):1459–1475. doi:10.1007/s00198-014-3012-y
179. Shin SH, Chae HD, Suprana A, et al. UTE MRI technical developments and applications in osteoporosis: a review. *Front Endocrinol*. 2025;16:1510010. doi:10.3389/fendo.2025.1510010
180. Zhao Y, Yan J, Zhu Y, et al. A novel prognostic 6-gene signature for osteoporosis. *Front Endocrinol*. 2022;13:968397. doi:10.3389/fendo.2022.968397
181. Ding H, Meng J, Zhang W, et al. Medical examination powers miR-194-5p as a biomarker for postmenopausal osteoporosis. *Sci Rep*. 2017;7(1):16726. doi:10.1038/s41598-017-17075-w
182. Lin B, Pan Z. Consensus gene modules related to levels of bone mineral density (BMD) among smokers and nonsmokers. *Bioengineered*. 2021;12(2):10134–10146. doi:10.1080/21655979.2021.2000746
183. Yang W, Xia S, Jia X, et al. Utilizing surface-enhanced Raman spectroscopy for the adjunctive diagnosis of osteoporosis. *Eur J Med Res*. 2024;29(1):476. doi:10.1186/s40001-024-02081-2
184. Liang Q, Zhao J, Yang F, et al. Face2Bone explainable AI model predicts osteoporosis risk from facial images in proof of concept study. *Sci Rep*. 2025;15(1):40913. doi:10.1038/s41598-025-20462-3
185. Bersanelli M, Mosca E, Remondini D, et al. Methods for the integration of multi-omics data: mathematical aspects. *BMC Bioinf*. 2016;17(2):S15. doi:10.1186/s12859-015-0857-9
186. Keenan L, Lorna Younger R, Race PR, et al. Molecular life sciences in the era of the fourth industrial revolution: sequencing, multi-omics and artificial intelligence. *Emerg Top Life Sci*. 2025;9(2):BSR20253294. doi:10.1042/ETLS20253019
187. Yang T, Xiao Y, Bao Z, et al. The rise and potential opportunities of large language model agents in bioinformatics and biomedicine. *Brief Bioinform*. 2025;26(6):bbaf601. doi:10.1093/bib/bbaf601
188. Zubair M, Khan AH, Bilal SF, et al. Deep learning approaches for resolving genomic discrepancies in cancer: a systematic review and clinical perspective. *Brief Bioinform*. 2025;26(6):bbaf541. doi:10.1093/bib/bbaf541
189. Du J, Chang M, Jiang K, et al. Identification of therapeutic targets for osteoporosis through multiomics integration and the potential dual role of androgen receptor signaling in its pathophysiology. *J Chem Inf Model*. 2025;65(16):8833–8845. doi:10.1021/acs.jcim.5c00386
190. Ziaastani Z, Kalantari-Khandani B, Niazi M-J, et al. Identification of critical genes and metabolic pathways in rheumatoid arthritis and osteoporosis toward drug repurposing. *Comput Biol Med*. 2024;180:108912. doi:10.1016/j.compbiomed.2024.108912
191. Zhang L, Yang Y, Geng D, et al. Identification of potential therapeutic targets and molecular regulatory mechanisms for osteoporosis by bioinformatics methods. *BioMed Res Int*. 2021;2021(1):8851421. doi:10.1155/2021/8851421

192. Guan Y, Ackert-Bicknell CL, Kell B, et al. Functional genomics complements quantitative genetics in identifying disease-gene associations. *PLoS Comput Biol*. 2010;6(11):e1000991. doi:10.1371/journal.pcbi.1000991
193. Feng Z-W, Xiao H-F, Wang X-W, et al. Unraveling key m6A modification regulators signatures in postmenopausal osteoporosis through bioinformatics and experimental verification. *Orthop Surg*. 2024;16(6):1418–1433. doi:10.1111/os.14064
194. Yang C, Ren J, Li B, et al. Identification of gene biomarkers in patients with postmenopausal osteoporosis. *Mol Med Rep*. 2019;19(2):1065–1073.
195. Long S-W, Li S-H, Li J, et al. Identification of osteoporosis ferroptosis-related markers and potential therapeutic compounds based on bioinformatics methods and molecular docking technology. *BMC Med Genomics*. 2024;17(1):99.
196. Xiao K-W, Li J-L, Zeng Z-H, et al. Monocytes affect bone mineral density in pre- and postmenopausal women through ribonucleoprotein complex biogenesis by integrative bioinformatics analysis. *Sci Rep*. 2019;9(1):17290. doi:10.1038/s41598-019-53843-6
197. Hao S, Xinqi M, Weicheng X, et al. Identification of key immune genes of osteoporosis based on bioinformatics and machine learning. *Front Endocrinol*. 2023;14:1118886. doi:10.3389/fendo.2023.1118886
198. Chen L, Zhao Y, Qiu J, et al. Analysis and validation of biomarkers of immune cell-related genes in postmenopausal osteoporosis: an observational study. *Medicine*. 2024;103(19):e38042. doi:10.1097/MD.00000000000038042
199. Zhang B, Pei Z, Tian A, et al. Multi-omics analysis to identify key immune genes for osteoporosis based on machine learning and single-cell analysis. *Orthop Surg*. 2024;16(11):2803–2820. doi:10.1111/os.14172
200. Li J, Li J, Zheng M, et al. Elucidating the role of FBXW4 in osteoporosis: integrating bioinformatics and machine learning for advanced insight. *BMC Pharmacol Toxicol*. 2025;26(1):20. doi:10.1186/s40360-025-00844-z
201. Wang X, Meng L, Zhang J, et al. Identification of ferroptosis-related molecular clusters and genes for diabetic osteoporosis based on the machine learning. *Front Endocrinol*. 2023;14:1189513. doi:10.3389/fendo.2023.1189513
202. Feng Z, Wu Z, Zhang Y. Integration of bioinformatics and machine learning approaches for the validation of pyrimidine metabolism-related genes and their implications in immunotherapy for osteoporosis. *BMC Musculoskelet Disord*. 2024;25(1):402. doi:10.1186/s12891-024-07512-z
203. Bi K, Chen Y, Hu Y, et al. Comprehensive bioinformatics analysis reveals novel potential biomarkers associated with aging and mitochondria in osteoporosis. *Sci Rep*. 2025;15(1):934. doi:10.1038/s41598-024-84926-8
204. Li S, Chen B, Chen H, et al. Analysis of potential genetic biomarkers and molecular mechanism of smoking-related postmenopausal osteoporosis using weighted gene co-expression network analysis and machine learning. *PLoS One*. 2021;16(9):e0257343. doi:10.1371/journal.pone.0257343
205. Zeng H-B, Yuan X-T, Wu L-G, et al. Exploring the role of mitochondrial dysfunction-related genes in osteoporosis using weighted gene co-expression network analysis and machine learning. *Eur J Med Res*. 2025;31(1):103. doi:10.1186/s40001-025-03675-0
206. Mishra BH, Mishra PP, Raitoharju E, et al. Modular genome-wide gene expression architecture shared by early traits of osteoporosis and atherosclerosis in the Young Finns Study. *Sci Rep*. 2021;11(1):7111. doi:10.1038/s41598-021-86536-0
207. Du A, Xu R, Yang Q, et al. Exploration of shared gene signatures and molecular mechanisms between type 2 diabetes and osteoporosis. *J Cell Mol Med*. 2024;28(9):e18141. doi:10.1111/jcmm.18141
208. Zhao R, Xiong C, Zhao Z, et al. Exploration of the shared hub genes and biological mechanism in osteoporosis and type 2 diabetes mellitus based on machine learning. *Biochem Genet*. 2023;61(6):2531–2547. doi:10.1007/s10528-023-10390-0
209. Liu J, Zhang D, Cao Y, et al. Screening of crosstalk and pyroptosis-related genes linking periodontitis and osteoporosis based on bioinformatics and machine learning. *Front Immunol*. 2022;13:955441. doi:10.3389/fimmu.2022.955441
210. Xu G, Zhang W, Yang J, et al. Identification of neutrophil extracellular traps and crosstalk genes linking inflammatory bowel disease and osteoporosis by integrated bioinformatics analysis and machine learning. *Sci Rep*. 2023;13(1):23054. doi:10.1038/s41598-023-50488-4
211. Tang H, Hu K, He Y, et al. Identification of potential biomarkers for osteoporosis and chronic kidney disease through bioinformatics and machine learning algorithm. *PLoS One*. 2026;21(5):e0348515. doi:10.1371/journal.pone.0348515
212. Wu X, Park S. A prediction model for osteoporosis risk using a machine-learning approach and its validation in a large cohort. *J Korean Med Sci*. 2023;38(21):e162. doi:10.3346/jkms.2023.38.e162
213. Zhang C, Wang F, Ke J, et al. Exploring the shared genetic characteristics of peri-implantitis and osteoporosis: a transcriptomic analysis perspective. *Int Dent J*. 2025;75(5):103865. doi:10.1016/j.identj.2025.103865
214. Lv Y, Du Y, Lin P, et al. Exploring the molecular intersections of osteoporosis and sarcopenia: an integrated bioinformatics and experimental validation. *Exp Gerontol*. 2026;214:113033. doi:10.1016/j.exger.2026.113033
215. Lin S, Zhang J, Jan C, et al. Impact of artificial intelligence on the availability, accessibility, acceptability and quality of ophthalmic disease screening services: a scoping review. *Br J Ophthalmol*. 2025;bjo–2024–325174.
216. Tun HM, Naing L, Malik OA, et al. Artificial Intelligence (AI)-based tools in the diagnosis and management of prostate cancer: a systematic review and meta-analysis. *Prostate Cancer Prostatic Dis*. 2025. doi:10.1038/s41391-025-01060-w
217. Doppalapudi S, Qiu RG, Badr Y. Lung cancer survival period prediction and understanding: deep learning approaches. *Int J Med Inf*. 2021;148:104371. doi:10.1016/j.ijmedinf.2020.104371
218. Smets J, Shevroja E, Hügler T, et al. Machine learning solutions for osteoporosis-a review. *J Bone Miner Res Off J Am Soc Bone Miner Res*. 2021;36(5):833–851. doi:10.1002/jbmr.4292
219. Yoo TK, Kim SK, Kim DW, et al. Osteoporosis risk prediction for bone mineral density assessment of postmenopausal women using machine learning. *Yonsei Med J*. 2013;54(6):1321–1330. doi:10.3349/ymj.2013.54.6.1321
220. Jin W, Xu L, Yue C, et al. Development and validation of explainable machine learning models for female Hip osteoporosis using electronic health records. *Int J Med Inf*. 2025;199:105889. doi:10.1016/j.ijmedinf.2025.105889
221. de Vries BCS, Hegeman JH, Nijmeijer W, et al. Comparing three machine learning approaches to design a risk assessment tool for future fractures: predicting a subsequent major osteoporotic fracture in fracture patients with osteopenia and osteoporosis. *Osteoporos Int J Establ Result Coop Eur Found Osteoporos Natl Osteoporos Found USA*. 2021;32(3):437–449. doi:10.1007/s00198-020-05735-z
222. Park HW, Jung H, Back KY, et al. Application of machine learning to identify clinically meaningful risk group for osteoporosis in individuals under the recommended age for dual-energy X-Ray absorptiometry. *Calcif Tissue Int*. 2021;109(6):645–655. doi:10.1007/s00223-021-00880-x
223. Peng Y, Zhang C, Zhou B. A cross-sectional study comparing machine learning and logistic regression techniques for predicting osteoporosis in a group at high risk of cardiovascular disease among old adults. *BMC Geriatr*. 2025;25(1):209. doi:10.1186/s12877-025-05840-w

224. Hsu C-T, Huang C-Y, Chen C-H, et al. Machine learning models to predict osteoporosis in patients with chronic kidney disease stage 3-5 and end-stage kidney disease. *Sci Rep.* **2025**;15(1):11391. doi:10.1038/s41598-025-95928-5
225. Wei Q, Liu Z, Chen X, et al. Development and validation of an explainable machine learning model for predicting osteoporosis in patients with type 2 diabetes mellitus. *Front Endocrinol.* **2025**;16:1611499. doi:10.3389/fendo.2025.1611499
226. Yu X, Liu W, Chen X, et al. Construction of a novel online calculator for prediction of osteoporosis risk in Chinese type 2 diabetes patients. *Exp Gerontol.* **2025**;208:112819. doi:10.1016/j.exger.2025.112819
227. Klemm C, Yeo I, Cohen-Levy WB, et al. Artificial neural networks can predict early failure of cementless total hip arthroplasty in patients with osteoporosis. *J Am Acad Orthop Surg.* **2022**;30(10):467–475. doi:10.5435/JAAOS-D-21-00775
228. Liu L, Si M, Ma H, et al. A hierarchical opportunistic screening model for osteoporosis using machine learning applied to clinical data and CT images. *BMC Bioinf.* **2022**;23(1):63. doi:10.1186/s12859-022-04596-z
229. Mateo J, Usategui-Martín R, Torres AM, et al. Improving prediction of fragility fractures in postmenopausal women using random forest. *Comput Biol Med.* **2025**;196(Pt A):110666. doi:10.1016/j.combiomed.2025.110666
230. Sebro R, De la Garza-Ramos C. Machine learning for the prediction of osteopenia/osteoporosis using the CT attenuation of multiple osseous sites from chest CT. *Eur J Radiol.* **2022**;155:110474. doi:10.1016/j.ejrad.2022.110474
231. Sebro R, De la Garza-Ramos C. Can we screen opportunistically for low bone mineral density using CT scans of the shoulder and artificial intelligence? *Br J Radiol.* **2024**;97(1160):1450–1460. doi:10.1093/bjr/tqae109
232. Huang C-B, Hu J-S, Tan K, et al. Application of machine learning model to predict osteoporosis based on abdominal computed tomography images of the psoas muscle: a retrospective study. *BMC Geriatr.* **2022**;22(1):796. doi:10.1186/s12877-022-03502-9
233. Kang SJ, Kim JOR, Kim MJ, et al. Preventive machine learning models incorporating health checkup data and hair mineral analysis for low bone mass identification. *Sci Rep.* **2024**;14(1):18792.
234. Huang W, Liu D, Liu G, et al. Association of urinary heavy metals with osteoporosis in US adults using interpretable machine learning. *Toxicol Lett.* **2026**;417:111853. doi:10.1016/j.toxlet.2026.111853
235. Queralto JM, Torres J, Guinot M. Neural networks for the biochemical prediction of bone mass loss. *Clin Chem Lab Med.* **1999**;37(8):831–838. doi:10.1515/CCLM.1999.125
236. Sadatsafavi M, Moayyeri A, Soltani A, et al. Artificial neural networks in prediction of bone density among post-menopausal women. *J Endocrinol Invest.* **2005**;28(5):425–431. doi:10.1007/BF03347223
237. Jiang H, Guo J, Li J, et al. Artificial neural network modeling to predict neonatal metabolic bone disease in the prenatal and postnatal periods. *JAMA Network Open.* **2023**;6(1):e2251849. doi:10.1001/jamanetworkopen.2022.51849
238. Suh B, Yu H, Kim H, et al. Interpretable Deep-learning approaches for osteoporosis risk screening and individualized feature analysis using large population-based data: model development and performance evaluation. *J Med Internet Res.* **2023**;25:e40179. doi:10.2196/40179
239. Hung W-C, Lin Y-L, Cheng T-T, et al. Establish and validate the reliability of predictive models in bone mineral density by deep learning as examination tool for women. *Osteoporos Int J Establ Result Coop Eur Found Osteoporos Natl Osteoporos Found USA.* **2024**;35(1):129–141. doi:10.1007/s00198-023-06913-5
240. Cho SW, Hong N, Kim KM, et al. Spine age estimation using deep learning in lateral spine radiographs and DXA VFA to predict incident fracture and mortality. *Npj Aging.* **2025**;11(1):83. doi:10.1038/s41514-025-00271-8
241. Tang J, Yin X, Lai J, et al. Fusion of X-Ray images and clinical data for a multimodal deep learning prediction model of osteoporosis: algorithm development and validation study. *JMIR Med Inform.* **2025**;13:e70738. doi:10.2196/70738
242. Lin Y-T, Chu C-Y, Hung K-S, et al. Can machine learning predict pharmacotherapy outcomes? An application study in osteoporosis. *Comput Methods Programs Biomed.* **2022**;225:107028. doi:10.1016/j.cmpb.2022.107028
243. Li Q, Han X-C, Zhou S-R, et al. Discovery of novel cathepsin K inhibitors for osteoporosis treatment using a deep learning-based strategy. *Expert Opin Drug Discov.* **2025**;20(10):1345–1356. doi:10.1080/17460441.2025.2527686
244. Bonaccorsi G, Giganti M, Nitsenko M, et al. Predicting treatment recommendations in postmenopausal osteoporosis. *J Biomed Inform.* **2021**;118:103780. doi:10.1016/j.jbi.2021.103780
245. Sugawara Y, Shimizu T, Ishizu H, et al. Development of an explainable machine learning model to reproduce and interpret expert pharmacological decisions in osteoporosis treatment. *Bone.* **2026**;204:117745. doi:10.1016/j.bone.2025.117745
246. Fasihi L, Tartibian B, Eslami R, et al. Artificial intelligence used to diagnose osteoporosis from risk factors in clinical data and proposing sports protocols. *Sci Rep.* **2022**;12(1):18330. doi:10.1038/s41598-022-23184-y
247. Hong N, Whittier DE, Glüer C-C, et al. The potential role for artificial intelligence in fracture risk prediction. *Lancet Diabetes Endocrinol.* **2024**;12(8):596–600. doi:10.1016/S2213-8587(24)00153-0
248. Wang X, Chen J, Zheng Q, et al. Artificial intelligence in joint face-skeletal prediction for orthognathic procedures: a systematic review. *J Cranio-Maxillo-Fac Surg off Publ Eur Assoc Cranio-Maxillo-Fac Surg.* **2025**;53(12):2301–2309. doi:10.1016/j.jcms.2025.11.002
249. Bai L, Xia Z, Triffitt JT, et al. Generation artificial intelligence (GenAI) and Biomaterials Translational: steering innovation without misdirection. *Biomater Transl.* **2024**;5(1):1–2. doi:10.12336/biomatertransl.2024.01.001
250. Jiang Y, Salley D, Sharma A, et al. An artificial intelligence enabled chemical synthesis robot for exploration and optimization of nanomaterials. *Sci Adv.* **2022**;8(40):eabo2626. doi:10.1126/sciadv.abo2626
251. Qiu S, Ji P, Wang Y, et al. Osteoimmune-regulative metal-organic framework nanomedicine for effective osteoporosis therapeutics. *Biomaterials.* **2026**;333:124184. doi:10.1016/j.biomaterials.2026.124184
252. Prakashan D, Pr R, Kaushik A, et al. Sustainable nanotechnology and artificial intelligence to empower image-guided therapy for precision healthcare. *BME Front.* **2025**;6:0150. doi:10.34133/bmef.0150
253. Li X, Li G, Zhao Y, et al. Exploring and comparing the use of large language models in supporting osteoporosis health consultations. *Clin Interv Aging.* **2025**;20:2133–2143. doi:10.2147/CIA.S551572
254. Querrer R, Vieira LS, Teodoro AB, et al. Deep learning for osteoporosis screening in dental practice: a systematic review. *Dento Maxillo Facial Radiol.* **2026**;55(1):1–27. doi:10.1093/dmfr/twaf052
255. Zhao R, Yang H, Li Y, et al. The diagnostic value of image-based machine learning for osteoporosis: systematic review and meta-analysis. *J Med Internet Res.* **2026**;28:e75965. doi:10.2196/75965

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